

SPECTROPHOTOMETER

OPERATING INSTRUCTIONS



BECKMAN

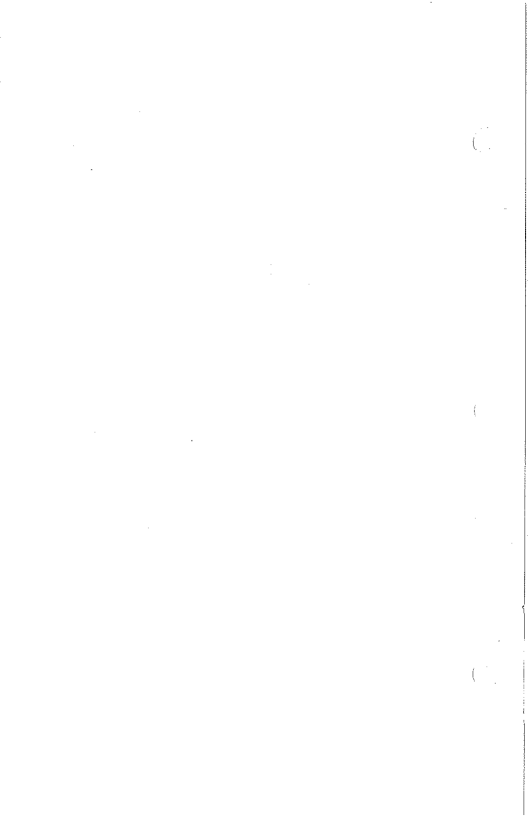
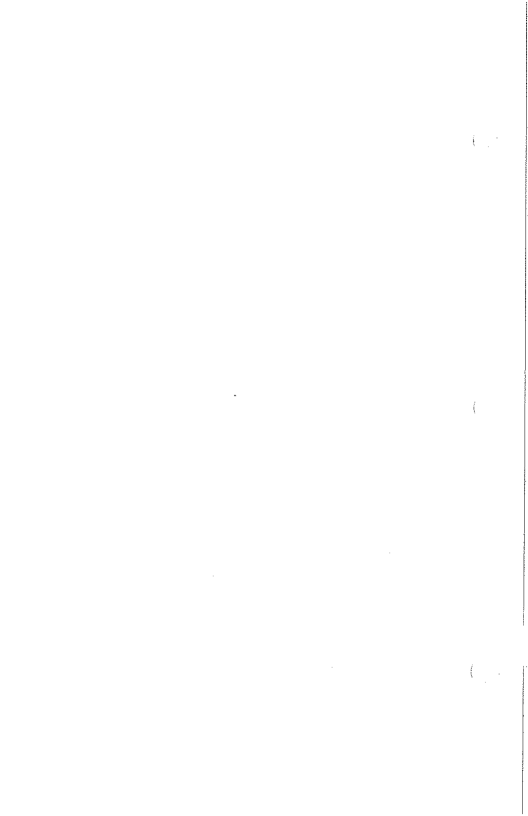


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SECTION ONE

INTRODUCTION

1.1 General Description

The DU® Series 600 Spectrophotometer is a microprocessor controlled spectrophotometer intended for use in quantitative and qualitative biological research and industrial procedures that require spectrophotometric measurements in the UV-visible region of the electromagnetic spectrum. If the instrument is used in a manner other than as described, the safety and performance of the instrument can be impaired.

The DU Series 600 Spectrophotometer operates in the wavelength range of 190 to 1100 nm. Models of the instrument are available with either a monochrome or a color video display. Data storage on a 3½ inch diskette is optional. Various optional accessories are available, to configure the instrument for specific application needs.

The instrument features a graphic video display, which provides operational information using windowing techniques, and a "mouse" for operator control. The mouse is used to position an arrow on the window. When the arrow points to the desired position, the left button on the mouse is pressed to initiate the desired action. In these instructions, the positioning of the arrow and pressing the left mouse button is called "clicking on".

The instrument has two rapid reading modes: RediRead for taking readings at a fixed wavelength and RediScan for making a wavelength scan.

The instrument has three standard Routine Measurement modes. They include:

Fixed Wavelength - Takes absorbance or transmittance readings at up to 12 wavelengths. Readings at each wavelength can be multiplied by a factor.

Wavelength Scan - Performs wavelength scans in absorbance or transmittance. Data are automatically stored for manipulations including Trace, zoom, overlay and tabulate. Calculations include peak pick,

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valley pick, point pick, first to fourth derivative, log of absorbance, scatter correction, spectral addition, subtraction and multiplication, and net absorbance. Repetitive scanning is also performed in this mode.

Kinetics/Time - Calculates the rate of an absorbance versus time reaction with a choice of blank subtraction and graphic display of the data for multiple samples. Data are automatically stored for manipulations including Trace, zoom, overlay and tabulate.

Other Application modes are also available, which include:

Protein Analysis - Calculates the protein concentration using the Bradford, Lowry, Biuret, direct UV method, colloidal gold, and bicinchoninate (BCA). Prepares a standard curve using up to 30 standards. The user can choose to add, delete or rerun individual standards based upon a statistical analysis of the standard curve.

Nucleic Acid - Determines protein impurity in nucleic acid samples based upon the ratio of readings at two wavelengths with a choice of background correction. Protein and nucleic acid concentrations can also be calculated using the Warburg and Christian¹ coefficients.

Fraction Read/Plot - Collects, plots and tabulates data from a set of related fractions. Readings can be corrected for dilution. After data collection, individual fractions can be added, deleted or rerun. Data are plotted versus either fraction number or volume. Other data obtained for the fractions can be input and plotted with the absorbance data.

Single component quantitative analysis - Calculates a standard curve from up to 30 standards using either linear or nonlinear least-squares regression. Performs statistical analysis on the standard data. Allows the user to add, delete or rerun individual standards to optimize the calibration. Calculates the concentration of samples from the calibration data.

Enzyme Mechanism - Guides the operator through the necessary steps to calculate Km and Vmax. Displays the following plot types: Michaelis-Menten, Lineweaver-Burk, Eadie-Hofstee, and Hanes-Woolf. Calculates the Hill constant from the Hill Plot. Graphs inhibitor plots to determine Ki.

¹Warburg, O. and Christian, W., <u>Biochem. Z</u>:310, p. 384f (1942).

Enzyme Activity - Calculates the enzyme activity of large numbers of samples.

Multicomponent analysis - Uses Full Spectrum Quantitation (FSQTM) to calculate the concentration of up to ten components in a mixture from up to 32 standards which are mixtures. Performs a statistical analysis of the standard data to determine the accuracy of the calibration. Allows individual standards to be added, deleted or rerun to optimize the calibration. Calculates the concentration of samples from the calibration data.

Gel Scan - Collects and plots absorbance data as a function of distance for a sample prepared by electrophoresis. Calculates peak and valley locations, which are used for subsequent area calculation or molecular weight determination.

Performance Validation - Provides a simple procedure to verify the performance of an instrument. Tests which are performed include: wavelength accuracy and repeatability, resolution, baseline flatness, noise and stability.

One model of the instrument also contains a program mode, which allows the user to customize applications by writing programs which blank, collect and store data at the desired wavelengths, calculate results from the data, prompt the operator, format and label the output and control sampling accessories.

The instrument is provided with a parallel output for a Dot Matrix Printer and a bidirectional RS-232 communications port that can be configured for either communications or an X-Y Plotter. The communications port can be used for data transfer to and from the instrument and for remote control of the instrument by an external computer.

A full line of modular sampling accessories is available for the instrument. Included are temperature controlled cell holders, automatic multi-position cell holders, a batch sampler, sipper samplers, and long pathlength cell holders. The automated sampling accessories (sipper samplers, batch sampler, and auto samplers) are designed to operate in the modes described previously to automate analyses. Microsampling capabilities include the use of the 50 μL Microcell, the 100 μL Multi-Microcell, and the 5 μL Ultra-Microcell.

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1.2 Principles of Operation

Optical Principle

The optical diagram of the DU Series 600 Spectrophotometer is shown in Figure 1-1.

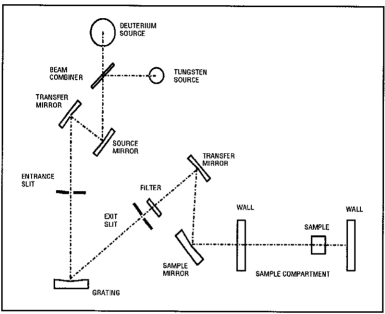


Figure 1-1.

DU Series 600 Spectrophotometer Optical Diagram

The DU Series 600 Spectrophotometer is a single beam spectrophotometer. Light from both sources enters the monochromator, where it is dispersed by the concave holographic grating. Monochromatic light exits the monochromator and illuminates the sample. The amount of light that passes through the sample is measured by the single photodiode detector.

The focal point of the beam in the sample compartment is on the right-hand side. This location permits the maximum amount of transmitted light to reach the detector from scattering samples. All sampling accessories position the sample at the focal point for best performance with microsamples and gels.

Blanking Method

A blank is always required before data collection; any reading taken without a blank is invalid. A blank reading is taken when <<BLANK>> (located in the permanent menu bar on the bottom of the window) is clicked on.

NOTICE

In the RediRead Mode the blank command is < ReadBlank >. In the RediScan Mode, the blank command is < ScanBlank >.

When the instrument blanks, the following steps are performed:

- 1. The monochromator is moved to the proper wavelength. This is the specified wavelength for a single wavelength reading.
- 2. The proper detector gain value is selected automatically. This minimizes the noise level and maximizes photometric accuracy.
- 3. Dark current is measured and corrected. This compensation assures accurate readings at high absorbance.
- 4. In the Wavelength Scan mode, only, a background scan is made. The blank (or reference) is automatically scanned over the same range at the same speed that the samples will be scanned, so that the background correction is optimal.

This calibration assures repeatable readings every time the instrument is used.

In all modes, a blank solution should be in the sample compartment during the blank. It is suggested that the solvent used to prepare the samples be used for the blank. However, air (no sample) may be used. A new blank reading should be taken each time the solvent is changed.

NOTICE

Plastic cuvettes, glass (Pyrex) cuvettes, and some solvents have significant absorption in the UV region. Verify that they transmit UV light by scanning them versus air before using them in the UV region.

To re-zero the instrument at any time between samples, insert the same blank solution and click on << BLANK>>.

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The instrument stores the blank and uses it until either the sources are turned off or another blank reading is taken. For best results, the instrument should be blanked frequently, allowing the blank reading to be taken shortly before the sample measurement is taken. A new blank should be read if the instrument has not been used for an hour.

Scanning

The background scan, made as part of the Blank procedure, is stored in the instrument and can be reused for an unlimited number of sample scans as long as the range and scan speed remain the same; unless the user blanks, the sources are turned off or the instrument is turned off. (The range can be decreased as long as the scan speed remains constant and no new background is required.) When a new background is required, it is indicated on the display.

A new background scan should be made every time a solvent is changed, because the background spectrum will likely be different. A new background scan should also be made if no scan has been made for over an hour. To rescan the background, click on <<BLANK>>, while in the Wavelength Scan mode.

The selected scanning speed determines the distance between each data point that is collected as the instrument scans through the chosen region. At 1200 nm/min, a data point is collected every nanometer. At 600 nm/min, a data point is collected every half nanometer.

As the sample data are collected the background is subtracted and the difference in absorbance (or ratio in transmittance) is plotted on the display.

Read Average Time

The noise level of the instrument, and therefore the uncertainty of a sample reading, is decreased by taking a number of readings and averaging them. The instrument takes a reading every 0.05 second. It takes a series of these readings over a user-selected time and averages them to obtain the blank and sample readings. For example, with a read average time of 0.5 seconds, ten readings are taken and averaged. The operator can specify a read average time from 0.05 to 99.9 seconds in all modes except Wavelength Scan and Multicomponent Analysis.

Read averaging is not used in the RediScan and Wavelength Scan modes. Background and sample scans are collected without averaging. Smoothing is used to improve the appearance of the data.

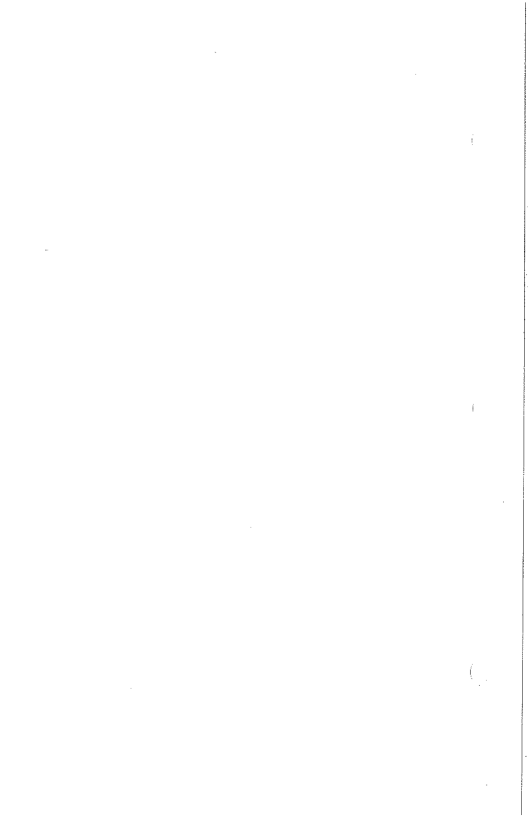
Smoothing

The displayed wavelength scan can be smoothed using a selectable smoothing function. The calculation, using the Savitzky and Golay² coefficients (as modified for end points by Peter A. Gorry³), is done for every data point in the scan, using the data points before and after the point of interest. The user selects the total number of data points used for the calculation, from 7 to 25. Use of too few points does not rid the scan of noise. Use of too many points can cause real peaks to be combined.

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²Savitzky, A., and Golay, M., Anal Chem, 36, 1964, p1627f.

³Gorry, Peter A., Anal Chem, 62, 1990, p570f.



2.1 Installation Instructions

The DU Series 600 Spectrophotometer is user installable. As an option, it can be installed by a qualified Beckman Field Service Engineer. The instructions for user installation are provided in Manual 517315.

Location

The DU Series 600 Spectrophotometer is designed to sit on a lab bench or table, which is level and flat and is capable of supporting its weight and the weight of all accessories.

The instrument is designed to operate in a clean laboratory environment, free from dust, fumes, excessive moisture, and corrosive chemicals. It should not be exposed to drafts from heating and cooling vents, heating elements, open windows or doors. Lab areas that receive direct sunlight should also be avoided.

An ambient room temperature of 15 - 40°C (59 - 104°F) should be maintained. Relative humidity should be 85% or less.

Instrument performance can be affected by strong electromagnetic fields that can exist in the proximity of large electric motors, centrifuges, diathermy machines and microwave sources.

The batch sampler <u>must</u> be placed adjacent to the right-hand side of the instrument. The other accessories can be placed in a convenient place, near the instrument, and within reach of the cables. Additional space is required for air circulation around the instrument and accessories for proper performance. Do not block these air spaces.

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2.2 Utility Requirements

Each of the following needs its own electrical outlet: the spectrophotometer, display, dot matrix printer, X-Y Plotter, batch sampler, and Peltier temperature controller. The electrical requirements of each are summarized in Table 1 for 100/120V and Table 2 for 220/240V systems.

The same circuit should be used for the instrument and all accessories. A dedicated circuit is preferred. Do not use a circuit which is also used by equipment that operates intermittently and creates wide fluctuations in power demand, such as refrigerators, water baths and centrifuges.

<u>For 100/120V operation</u>. The power line should provide three-wire single phase power. To provide multiple outlets, grounding type power strips may be used. Extension cords or multiple outlet adapters should not be used.

<u>For 220/240V operation</u>. The power line should provide three-wire single phase power. To provide multiple outlets, grounding type power strips may be used. Extension cords or multiple outlet adapters should not be used.

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	Frequency (Hz)	Voltage (VAC)	Current (Amps)
Spectrophotometer	50/60	100/120V±10%	1.8
Display	50/60	110V±10%	0.8
Dot Matrix Printer	50/60	120V±10%	0.6
X-Y Plotter	48 - 66	120V -10%,+5%	0.2
Batch Sampler	50/60	117V±10%	0.8
Peltier Temperature Controller	50/60	120V±10%	1.0

Table 1. Electrical Requirements for 100/120V Systems

	Frequency (Hz)	Voltage (VAC)	Current (Amps)
Spectrophotometer	50/60	220/240V±10%	0.9
Display	50/60	220V±10%	0.4
Dot Matrix Printer	50/60	220/240V±10%	0.3
X-Y Plotter	48 - 66	240V -10%,+5%	0.1
Batch Sampler	50/60	234V±10%	0.4
Peltier Temperature Controller	50/60	220/240V±10%	0.5

Table 2. Electrical Requirements for 220/240V Systems

2.3 Power Up

The DU Series 600 Spectrophotometer is powered up using the following steps.

1. Verify that the voltage on the rating plates on the instrument, display and printer are the same as the power source.

A CAUTION

The voltage indicated on the display, the instrument and the printer must be the same as the power source. If the incorrect voltage is indicated, do not plug in the instrument and contact the local Beckman service office.

The instrument, display and printer must be plugged into grounded power outlets.

NOTICE

If a power cord is not supplied, contact the local Beckman service office.

- 2. Plug the power cords on the instrument, the display and the printer into grounded, three-prong outlets that are on the same power line.
- Turn on the display. The indicator light on the front of the display should illuminate.
- 4. Turn on the instrument. The power switch is located on the right-hand side of the back of the instrument. The instrument fan should turn on.

If the fan does not turn on, turn off the instrument and check the fuses. Directions for fuse replacement are provided in the Corrective Maintenance section of this manual.

5. The display should illuminate, a message window with "Executing Power Up Diagnostics" should be displayed, and an arrow should appear on the display. The arrow should move when the mouse is moved.

If the display does not illuminate:

a. Verify that the indicator light on the display is illuminated, showing that there is power to the display.

- b. Verify that the brightness control on the display is adjusted properly.
- c. Verify that the cable on the display is connected to the "DISPLAY" port on the back of the instrument.

If the mouse does not move the arrow, verify that the cable attached to the mouse is connected to the "MOUSE" port on the back of the instrument.

NOTICE

If the recommended action does not correct the problem, power down the instrument, then power it up again. If the problem persists, contact the local Beckman service office.

- 6. Turn on the printer using the switch on the left-hand side of the printer. Verify that the "ON LINE" light is illuminated (or blinking).
- 7. Verify that the printer paper is loaded and that the top of the first page is aligned properly.

NOTICE

The instructions for loading of the paper and the replacement of the ribbon are located in Manual 514521.

2.4 Configuration

After the DU Series 600 Spectrophotometer has been installed and powered up, the Configuration mode is used to select parameters which setup the instrument, the output devices, and the sampling accessories. It is also used to assign the passwords for method protection. Configuration parameters are generally selected when the instrument is installed and are not changed during operation of the instrument.

The DU Series 600 Spectrophotometer user interface operates on the principle of windows. The "mouse" is used to position an arrow on the window. When the arrow points to the desired position, the left button on the mouse is pressed to initiate the desired action. In these instructions, the positioning of the arrow and pressing the left mouse button is called "clicking on". If the Power Up Diagnostic window is displayed, click on <Quit> to remove the window and display the Main window.

The Configuration window, Figure 2-1, is used to select one of seven windows that are used for different configurations. It is displayed when "CONFIGURATION" is clicked on from the Main window.

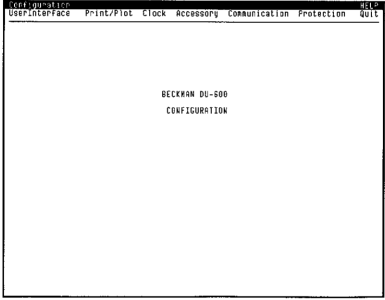


Figure 2-1. Configuration Window

NOTICE

Instructions to configure the instrument and Printer/Plotter, only, are provided in this section. Instructions for the X-Y Plotter are provided in Manual 514523. Instructions for the sampling devices are provided in Manual 517314.

User Interface Configuration

The User Interface Configuration window, Figure 2-2, is displayed when <userInterface> is clicked on from the Configuration window.

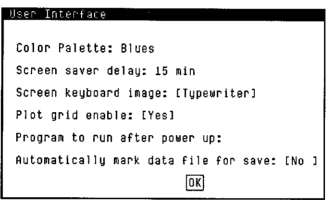


Figure 2-2. User Interface Configuration Window

Set the following:

- 1. Color palette This determines the set of colors used on the display for a color monitor, or if a monochrome monitor is used. Click on the current selection to display a window with the palette options. Click on the box before the palette of choice, then [OK] to remove the window.
- 2. Screen saver delay (minutes) The instrument dims the display whenever the instrument is not used for the selected amount of time and both sources are turned off. Click on and input the desired amount of time.
- 3. Screen keyboard image This determines the order of keys on the alphanumeric keypad. Click on to toggle between [Typewriter] and [Alphabetic].
- 4. Plot grid enable This determines whether a grid is displayed on graphs. Click on to toggle between [Yes] and [No] to enable or disable the grid, respectively.

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The input to this selection determines whether the grid is plotted on printouts generated on the Dot Matrix Printer. For printouts on the X-Y Plotter, however, the grid is enabled or disabled on the Printer and Plotter Configuration window.

5. Program to run after power up - This allows a user-written program to be executed automatically after power up is completed. The program could turn on the sources, have a delay to allow the sources to warm up, then execute the Performance Validation tests. (The Program mode, which is used to write the program, is not included on all models.)

To select a program, click on the displayed program name to display the Program directory. Click on one of the program names to select it.

6. Automatically mark data file for save - This affects the Save Clear and Quit windows, only. If this is enabled, the box is automatically darkened for each data file when the window is displayed. If this is disabled, the box is darkened when it is clicked on or if the file name is input.

When the desired parameters are selected, click on [OK] to store the entries and remove the User Interface Configuration window from the display.

Clock Configuration

The Clock Configuration window, Figure 2-3, is displayed when <Clock> is clicked on from the Configuration window.

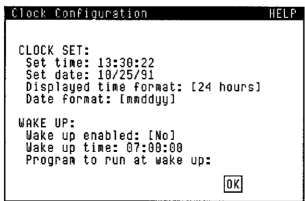


Figure 2-3. Clock Configuration Window

CLOCK SET

Input the following information to set the date and time, displayed in the permanent menu bar.

- 1. Set time Input the current time, using the 24 hour clock. The clock starts when [OK] is clicked on to remove the window.
- 2. Set date Input the current date.
- 3. **Time format** Select format for the time: [12 hours] or [24 hours]. If a 12 hour clock is selected, am and pm indications are *not* given.
- 4. **Date format** Select format for the date: [mmddyy], [ddmmyy], or [yymmdd].

WAKE UP

This allows the instrument to turn on the sources at an operator determined time, so that the instrument and the sources can be warmed up and ready to use. The instrument should be left with the Main window displayed and the sources turned off in preparation for Wake Up to execute.

- 5. Wake up enabled Toggle between [Yes] and [No] to enable or disable the wake up mode.
- 6. Wake up time Input the wake up time using the 24-hour clock.
- 7. Program to run at wake up This allows a user-written program to be executed automatically after the sources are turned on. The program could have a delay to allow the sources to warm up, followed by the Performance Validation mode or some other diagnostic tests. (The Program mode, which is used to write the program, is not included on all models.)

To select a program, click on the displayed program name to display the Program directory. Click on one of the program names to select it.

When the desired parameters are selected, click on [OK] to store the entries, start the clock and remove the Clock Configuration window from the display.

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Protection Configuration

The Protection Configuration window is used to identify each user and assign a password. The password prevents methods from being modified by another user. The master access code is required before passwords can be assigned or changed. This prevents unauthorized change of password.

To input or change passwords:

1. Click on **Protection>** from the Configuration window. The Protection Configuration window, Figure 2-4, is displayed.

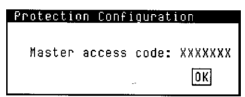


Figure 2-4.
Protection Configuration Window

2. Click on the X's that are displayed for the master access code, input the code, then click on [OK]. The Edit User Names and Passwords window, Figure 2-5, is displayed.

NOTICE

The master access code is assigned by Beckman Instruments and cannot be changed.

Edit	User	Names	and	Passwo	rds	
2. 3. 4. 5.	Tsafr Barru Mike Linda	Name k Aral rira Ma J Masta Simon a Whee rie Kol	eir er ian ler		1. 2. 34. 5. 6. 7. 8.	Password FA2 TM31 BSM0 MHS7 LW9 VK23
						OK

Figure 2-5.
Edit User Names and Passwords Window

- 3. Click on a position in the table in the "User Name" column and input a user name. Then click on the corresponding position in the "Password" column and input a password. Repeat for each user.
- 4. To edit any of the entries, click on it and input the desired information.
- 5. When all of the information has been input correctly, click on [OK] to store the information and remove the Edit User Names and Passwords window from the display. Then click on [OK] to remove the Protection Configuration window from the display.

Printer Configuration

The Printer and Plotter Configuration window, Figure 2-6, is displayed when **<Print/Plot>** is clicked on from the Configuration window.

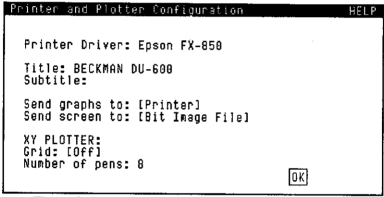


Figure 2-6. Printer and Plotter Configuration Window

Select the following parameters:

- 1. Printer Driver Selects the printer type being used. If a HP Desk Jet is selected, all text screens with 80 characters or less are printed portrait, and all graphs and all text screens with more than 80 characters are printed landscape, and, if any line has more than 100 characters, the print is condensed. To use the Cannon BJ-200, select the Epson LQ-510 and make sure that the DIP switches are set for the Epson LQ-510 mode.
- 2. Title Input up to 60 alphanumeric characters which are printed at the top of each printout made with the <Print> command. This can be used to identify the institution or department where the data were generated.

- 3. Subtitle Input up to 60 alphanumeric characters which are printed on the second line of each printout made with the <Print> command. This can be used to provide an analysis description or to identify the laboratory where the data were generated.
- 4. Send graph to If both the printer and plotter are installed, select from [Printer] and [XY Plotter]. This option determines where a printout, which includes a graph, is printed.
- 5. Send screen to This determines where a screen copy is sent when [PrtScrn] is clicked on. Select from either [Printer] or [Bit Image File]. If [Bit Image File] is selected, the information on the screen is made into a TIFF file. The TIFF file is named by the user and the file is placed in the ASCII directory.
- 6. Grid This parameter is used for the X-Y Plotter, only. Refer to Manual 514523 for more information.
- 7. Number of pens This parameter is used for the X-Y Plotter, only. Refer to Manual 514523 for more information.

When the desired parameters are input, click on [OK] to store the entries and remove the Printer and Plotter Configuration window from the display, then <Exit> to return to the Main window.

Installation

2.5 Calibration

There are three calibrations that are performed by the DU Series 600 Spectrophotometer: absorbance (or transmittance), wavelength and scan gain.

Absorbance (or transmittance) - The absorbance reading is calibrated each time a blank reading is taken. Blanking is discussed in section 1.2.

Wavelength - The wavelength is calibrated at the factory prior to shipment. At any time after installation, the wavelength can be recalibrated using the following instructions:

- 1. Verify that both sources are turned on. If either is off, turn it on by clicking on the appropriate command in the permanent menu bar.
- 2. With the Main window displayed, click on "DIAGNOSTICS" to display the Diagnostics window.
- Click on <Calibrate λ>. The instrument finds the deuterium emission line at 656.1 nm, recalibrates the wavelength and stores the information. (The instrument also checks the scan gain values and adjusts them, if necessary. Scan gain is discussed below.)
- 4. When the wavelength calibration is complete, click on <Quit> to remove the Diagnostics window and display the Main window.

Under normal operating conditions, it should not be necessary to recalibrate the wavelength on a routine basis. However, it may be desirable to recalibrate the wavelength after moving the instrument.

Scan Gain

The scan gain is set at the factory prior to shipment. The gain is checked and adjusted each time that the wavelength is calibrated, as described above. When a wavelength scan is performed using a blank that has significant absorbance in the scanning range, it may be desirable to change the scan gain settings in the instrument to compensate for the absorbance of the blank. This will increase the dynamic range of the instrument when scanning using the absorbing blank. To change the scan gain:

 With the Main window displayed, click on "DIAGNOSTICS" to display the Diagnostics window.

Installation 2-13

- 2. Place a cuvette of the blank in the cell holder.
- 3. Click on <SetScanGain>. The instrument adjusts the scan gain values to compensate for the absorbance of the blank.
- 4. When the scan gain has been set, click on <Quit> to remove the Diagnostics window and display the Main window.

NOTICE

The instrument will continue to use the new gain values until the gain is adjusted using either $\langle Calibrate \lambda \rangle$ or $\langle SetScanGain \rangle$. It may be desirable to repeat the above procedure with nothing in the sample compartment after changing the gain for a significantly absorbing blank.

2.6 When Not in Use

When the instrument is not in use for more than two hours, turn off both sources. The power to the instrument can, but does not need to, remain turned on. The "Screen Saver Delay" set in the User Interface Configuration Window, dims the display after the selected time. Any movement of the mouse will re-illuminate the display.

Before using the instrument again, turn on either or both sources, depending upon the application, and allow them to warm up for at least 30 minutes before taking readings.

GENERAL OPERATING INFORMATION

The DU Series 600 Spectrophotometer user interface operates on the principle of windows. A "mouse" is used to move an arrow around a window. Action is initiated when a mouse button is clicked on.

The instrument contains several analysis modes, which are selected from the Main window by using the mouse to position an arrow on the mode name and clicking on the left mouse button. When the mode is selected, an analysis window for the mode is displayed. Analysis parameters can be input in the analysis window or can be recalled from a stored method.

The analysis modes have an associated Method window, which is used to recall stored methods and to input the parameters to develop new methods. The methods, sample data, calculated results, and standard data can be stored, in a designated file. These files are located in the main instrument or in an optional external disk.

3.1 Analysis Windows

Analysis windows are displayed during normal operation. These windows are used to select parameters, collect data and calculate results. A typical analysis window is shown in Figure 3-1.

	Fired Wave ReadSample		h Meth	od Par	ameters	S	aveClear	Print	HEL⊃ Quit
>	Results f Read aver			1 0 Read mo	ode: [Abs]		name: A:\F g device:		•
·	Sample I	D	Fact	350.0 or 56.00	Facto	440.0 or 230.0	λ 5 Factor	20.0 6.500	
	1 2 43F 43T 46J 48K		Abs 0.2790 0.3647 0.6747 0.67413 1.0447 0.9504	RBSU 1 t RG/M1 15.6221 28.4213 37.7840 35.9108 58.5056 53.2240	Abs 0.1535 0.0971 0.2244 0.2421 0.3162 0.3767	Result ng/ml 35.2955 22.3246 51.6012 55.6869 72.7205 86.6482	Abs 6.3152 6.3784 1.0832 6.6864 1.4383 1.3868	Result ng/n1 2.6490 2.4596 7.8407 4.4613 9.2369 8.9698	
	BLANK Match off		OFF] Re			486.0 nm .0813 Abs		DATE TEMP /30/91 N/A	CELL 1

Figure 3-1. Typical Analysis Window

The manual uses the following nomenclature when referring to analysis windows:

1 - Window

The entire display, or any portion of the display that is enclosed with a box. More than one window can be displayed at a time.

2 - Window Name

All analysis windows have a name on the top line of the window. An example of each analysis window is provided in this manual. A listing of these windows is provided at the end of the Table of Contents.

3 - Help Messages

"HELP" is displayed on the top line of the window, to the right of the window name. Click on "HELP" to display Help windows.

4 - Permanent Menu Bar

A list of commands is always displayed at the bottom of the display. A permanent menu bar command is referred to in these instructions with the use of double angle brackets, i.e. << BLANK>>. Status information is provided on the right-hand side of the permanent menu bar.

5 - Window Menu Bar

A list of commands is located at the top of each window, directly below the window name. The specific commands change with each window. A window menu bar command is referred to in these instructions with the use of single angle brackets, i.e. <Print>. Commands that can be used are identified by color.

6 - File Names

The files where the method, standard data, and/or results data are stored are listed directly under the window menu bar.

7 - Parameters

The analysis parameters are listed near the top of a window. Parameters are referred to in these instructions with the use of quotes, i.e. "Wavelength". The values for parameters that can be changed are displayed in a different color than the parameter name.

8 - Data Column Labels

Some column labels can be changed, such as the units for calculated results. Labels that can be changed are displayed in the same color as parameters that can be changed.

9 - Scrolling Arrows

Click on the arrows to display data that is not displayed because of insufficient room. The window shown in Figure 3-1 has both up and down arrows and right and left arrows. Many windows have only up and down arrows.

3.2 General Operating Procedure

When the DU Series 600 Spectrophotometer is powered up, it performs a series of diagnostic tests. When these tests are completed satisfactorily, the Power Up Diagnostics window, Figure 3-2, is displayed.

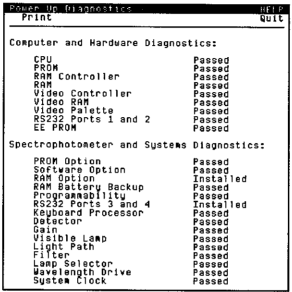


Figure 3-2. Power Up Window

If all power up tests pass, remove the Power Up Diagnostics window from the display by using the mouse to move the arrow to <Quit> and clicking on the left mouse button. The Main window, Figure 3-3, is displayed. The Main window is used to select an operating mode.

NOTICE

If any power up test fails, refer to the Troubleshooting section for instructions.

Programmability is an option and may not be installed on all instruments.

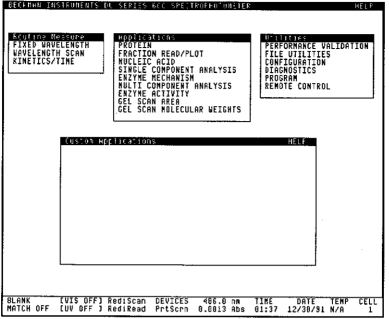


Figure 3-3. Main Window

The following are general operating instructions. Specific instructions for each analysis mode are provided in the respective section in this manual.

- To select an analysis mode from the Main window, use the mouse to move the arrow to the desired mode. Press the left mouse button to click on the analysis mode. An analysis window, such as shown in Figure 3-1, is displayed.
- Analysis parameters are listed on the window. The mouse is used to input different values for the parameters. More information on parameter input is given in section 3.7.
 - As an alternative, a stored method can be recalled or a new method can be developed. Either of these is done by clicking on <Method>, located on the menu bar at the top of the window to display the Method window. A typical Method window is shown in Figure 3-16.
- 3. When the desired parameters are displayed, place a cuvette of blank solution in the sample compartment and click on <<BLANK>>. (<<BLANK>> is located in the permanent menu bar.)

- 4. Place the first sample in the sample compartment and click on <Read-Samples>. (<ReadSamples> is located on the menu bar at the top of the window.) As an alternative, the right mouse button can be used to take a sample reading, with the cursor in any position.
- 5. To print the data, click on **<Print>**. All information in the window is printed, even if only partial information is displayed, because of insufficient room. If the window contains graphic data, it is printed on the device selected in the Printer and Plotter Configuration window. If the window contains no graphic data, it is printed on the printer, if installed and operational. The permanent menu bar is not printed.

To stop printing while the DU-600 is transferring the information to the printer, the Stop Printing window is placed at the top of the display. To stop printing, click on [QUIT]. Reset the paper to the top of a new page before starting another printout.

To stop printing after the Stop Print window is removed, click on <<DEVICES>> to display the Device Control window. Click on [STOP PRINTING], then <Exit> to remove the Device Control window from the display. Reset the paper to the top of a new page before starting another printout.

- 6. To clear all data from the window, with the option of storing the data, click on **SaveClear>**. If the data are stored, no additional data can be placed in the data file.
- 7. When the analysis is complete, click on <Quit> to return to the Main window. (<Quit> is located on the menu bar at the top of the window.) An opportunity is given to store the method and/or sample data.

1 -

3.3 Help Messages

Help windows are displayed by clicking on "HELP", displayed to the right of the window name. Typical Help windows are displayed in Figures 3-4 and 3-5. If a Help window similar to Figure 3-4 is displayed, click on the desired selection to display a Help window similar to Figure 3-5. Click on <Exit> to remove the Help windows from the display.

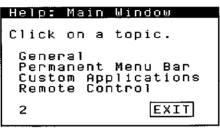


Figure 3-4. Help Window

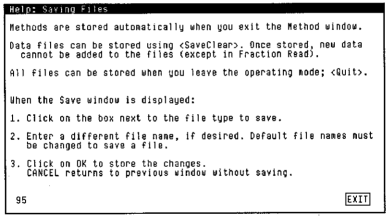


Figure 3-5. Help Window

3.4 The Mouse

The mouse, Figure 3-6, is used to move an arrow to desired locations on the DU Series 600 Spectrophotometer display. When the arrow is in a desired location, click on the left mouse button to initiate action. The items that can be clicked on include window menu bar commands, file names, parameters, sample data, and permanent menu bar commands.

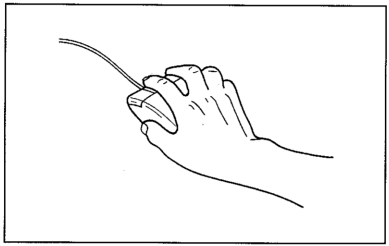


Figure 3-6. Mouse

The left mouse button is used the most frequently to click on a command or parameter. However, the mouse buttons have other uses:

- To take a reading The right mouse button can be used to initiate a reading, with the cursor in any position, as an alternate to <ReadSamples> in the window menu bar.
- 2. To input lower case letters When the alphanumeric keypad is displayed for character input, click on the right mouse button, rather than the left button, to input lower case letters.
- 3. To use Trace Trace is a feature that is used to find the ordinate and abscissa values from graphic data. Trace is a window menu bar command (<Trace>) that is clicked on in the normal manner. After <Trace> is clicked on, the mouse is used to move the arrow to the desired location on the graph and the center mouse button is clicked on to position a line on the graph. Then the right and left mouse buttons can be used to move the line to the right and left, respectively.

The Hour Glass

Most of the time the arrow appears on the window. However, when an action is initiated that cannot be completed quickly, the arrow is changed into an hour glass. The hour glass can be moved across the window in the same manner as the arrow, and the mouse buttons can be clicked on, but no action occurs until the arrow returns. Typical actions that cause the hour glass to be displayed include printing a window and performing complex data calculations.

The Diskette Symbol

When the optional disk drive is accessed, the arrow is changed into a diskette symbol. The diskette symbol cannot be moved and no action can be taken until disk access is completed.

3.5 Permanent Menu Bar

The permanent menu bar, Figure 3-7, is always displayed at the bottom of the display. Commands that can be used at any time during the operation of the instrument are listed on the left-hand side of the permanent menu bar. Current nanometer position and reading are displayed in the middle. Status information is displayed on the right-hand side. If a source burns out during operation, an error message is displayed under the commands on the left-hand side.

BLANK (VIS OFF) Rediscan DEVICES 486.0 nm TIME DATE TEMP CELL MATCH OFF (UV OFF) Rediread Priscrn 0.0013 Abs 81:37 12/30/91 M/A 1

Figure 3-7. Permanent Menu Bar

The permanent menu bar commands are:

BLANK

Take a reading at the analytical and background wavelength(s) and set the value 0.000 absorbance and 100%T. In the Wavelength Scan mode, only, make a background scan.

MATCH OFF/ON

If an Auto Cell Holder is used for the analysis, zero readings can be taken on each of the cuvettes. When Match is enabled, the zero readings are subtracted from all subsequent readings for the appropriate cell. Operational information is provided in Manual 517314.

VIS OFF/ON

Turn the visible source on or off.

UV OFF/WAIT/ON

Turn the UV source on or off. The UV source requires about 30 seconds to light after being turned on. During this time <<UV WAIT>> is displayed.

NOTICE

Do not blank the instrument or take sample readings while <<UV WAIT>> is displayed, even in the visible region. When the UV source lights, readings at all wavelengths are affected.

RediScan

Enter the RediScan mode. This is described in section 4.3.

RediRead

Enter the RediRead mode. This is described in section 4.2.

DEVICES

Display the Device Control window, Figure 3-8. The window is used to stop sending information to the printer or plotter after printing has begun; to position the transport at a cell position for the Auto Cell Holder, a millimeter position, or the home position to align; to control the temperature controller; and to move the aspirator arm on the batch sampler. When the desired action has been taken, click on **Exit>** to remove the window from the display.

The information input in the Device Control window is overridden if the device is automatically controlled as part of an analysis.

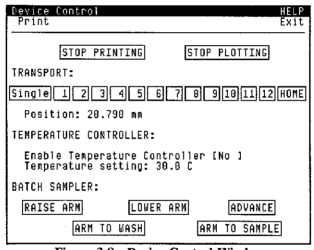


Figure 3-8. Device Control Window

PrtScrn

Print an exact copy of the entire display on the Dot Matrix Printer. Information that is not displayed because of insufficient room will not be printed. The X-Y Plotter cannot be used for this printout.

NOTICE

There is also a **Print>** command in the menu bar at the top of most windows, which is used to print the window in which it appears and also prints the continuation of data that are not displayed.

The status information includes:

Nanometer Position

The current nanometer position.

Absorbance/Transmittance Reading

The current reading, in either absorbance or transmittance, determined by the reading mode in the selected analysis mode. This field is blank until the instrument is blanked.

TIME

The current time. The time is set in the configuration mode, using the Clock Configuration window.

DATE

The current date. The date is set in the configuration mode, using the Clock Configuration window.

TEMP

The current temperature, if the Temperature Controller is being used. When the Temperature Controller is controlling temperature, "on" is displayed under the temperature. If the Temperature Controller is not installed or is not turned on, "N/A" is displayed. The Temperature Controller is turned on using the Device Control window, Figure 3-9.

CELL

If the Transport Accessory is installed, the Auto Cell Holder cell number that is in the beam is displayed. If the Transport Accessory is not installed, "N/A" is displayed.

3.6 Window Menu Bars

Across the top of analysis windows is a menu bar specific to the window. This is referred to in the manual as a window menu bar or simply a menu bar. The following are some of the commands that typically appear in the menu bar.

<ReadSamples>

Take a sample reading. In some modes, and with some sampling accessories, this command causes the window shown in Figure 3-9 to be displayed. Sample readings are started when [START] is clicked on and are stopped when [QUIT] is clicked on.

Insert sample(s) Click on START when ready Click on QUIT to stop START QUIT

Figure 3-9. Read Samples Window

In most cases, clicking on the right mouse button, with the cursor in any position, performs the same function as <ReadSamples> and [START].

<Window Name>

Most operating modes have more than one analysis window. The other windows are displayed by clicking on the window name, displayed in the menu bar.

<Method>

Display the Method window to select a stored method or create a new method.

<SaveClear>

Clear all data from the window, without leaving the analysis mode. Before the data are removed, the Save Clear window, Figure 3-10, is displayed, allowing the data to be stored. To store the data, click on the box to darken it, verify that the desired file name is displayed and click on [OK]. To change the file name, click on it to display the Results File Directory window, which is described in section 3.9. To change the storage location, click on the displayed location to toggle between [A:\] and [B:\].



Figure 3-10. Save Clear Window

NOTICE

The user can select to have the save box darkened automatically each time the Quit window is displayed. This selection is made on the User Interface window in the Configuration mode, which is described in section 2.4.

<Print>

Print the data displayed on the window. Alphanumeric data are printed on the Dot Matrix Printer. Sample data and other information that are not displayed on the screen because of insufficient room are printed. If the X-Y Plotter is installed, graphic data can be printed on either the Dot Matrix Printer or the X-Y Plotter. This is selected in the Printer and Plotter Configuration window.

<Exit>

Remove the window.

<Quit>

Leave an operating mode and return to the Main window. If the method, standards file or results file has changed, the Quit window, Figure 3-11, is displayed. To store any of the information, click on the associated box to darken it, verify that the desired file name is displayed and click on [OK]. To change a method file name, click on it to display the alphanumeric keypad and input the desired file name. To change a data file name, click on it to display the Results File Directory window, which is described in section 3.9. To change the storage location, click on the displayed location to toggle between [A:\] and [B:\].

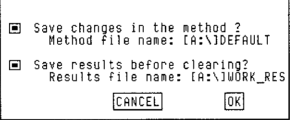


Figure 3-11. Quit Window

NOTICE

The user can select to have the save boxes darkened automatically each time the Quit window is displayed. This selection is made on the User Interface window in the Configuration mode, which is described in section 2.4.

3.7 Parameter Input

Parameters that pertain to a particular analysis window are listed near the top of the window. The Method window contains a complete listing of analysis parameters. The selected value can be changed for any parameter where the displayed value is a different color than the parameter. For example: the parameter, "Wavelength", is a different color that the value, "500".

There are several types of parameter input. These include:

Numeric Input

When a parameter is clicked on that requires a numeric input, the numeric keypad, Figure 3-12, is displayed. The limits of the input are displayed at the bottom of the keypad. Input the desired value and click on [OK] to accept the value and remove the numeric keypad. Examples of parameters that require numeric input include the wavelength, number of samples, and read average time.

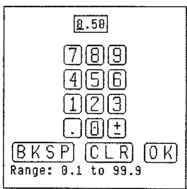


Figure 3-12. Numeric Keypad

As an alternative, the optional keyboard can be used to input numbers. When the numbers are input, they are displayed in the box at the top of the keypad. When the desired number(s) are displayed, press enter on the keyboard (or click on [OK] on the keypad) to accept the value and remove the keypad.

Alphanumeric Input

When a parameter is clicked on that requires an alphanumeric input, the alphanumeric keypad, Figure 3-13, is displayed. The maximum number of characters that can be input is indicated by the space provided for the input. Input the desired information and click on

[OK] to accept the information and remove the alphanumeric keypad. Examples of parameters that require alphanumeric input include file names, concentration units and sample identifications.

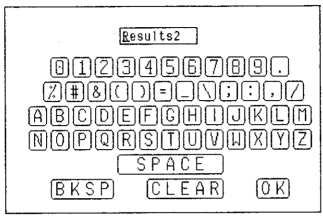


Figure 3-13. Alphanumeric Keypad

As an alternative, the optional keyboard can be used to input this information. When the characters are input, they are displayed in the box at the top of the keypad. When the desired characters are displayed, press enter on the keyboard (or click on [OK] on the keypad) to accept the information and remove the keypad.

Option Selection

If a parameter has a variety of options, a selection window is displayed when the parameter is clicked on. In some cases, the word "VIEW" is displayed and clicked on to get the selection window. A typical selection window is shown in Figure 3-14.

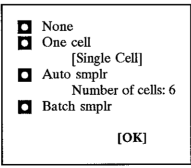


Figure 3-14.
Sampling Device Window

Each of the options on the selection window is preceded by a box. Click on the box to darken it to select the desired option. A second click on the box removes the darkened area. Some selection windows accept only one choice. Others allow multiple selections. When the selection(s) are made, click on [OK] to input the option and remove the selection window.

Bracketed Options

Some parameters require a selection from only two or three options, such as "yes" or "no". The selection for these types of parameters is displayed with brackets, such as [Yes]. When the selection in brackets is clicked on, the selection changes to the other option. For example, click on [Yes] to display [No]. Examples of parameters that have bracketed options include the selection of absorbance or transmittance readings and the use of a background wavelength.

Parameters that Cannot Change

If the selected value for an analysis parameter is displayed in the same color as the parameter, the value cannot be changed. There are several reasons that a parameter cannot be changed. They include:

- 1. The method in use has been protected. After a method is developed, it can be protected, so that the analysis must be done with the selected parameters.
- 2. Parameters used for sample collection must be the same as for the related standards. For example, if the standard curve was calculated using an analytical wavelength of 500 nm, the wavelength for the sample analysis must be 500 nm.
- 3. Data collection is complete. For example, the temperature cannot be changed if data were previously collected.

3.8 Method Development and Use

Parameter setup for analyses that are repeated often can be simplified by storing the analysis parameters in a method file. Each analysis mode (with the exception of the Fraction Read mode) has a Method window associated with it. The Method window is used to setup, store and retrieve analysis parameters. A typical Method window is shown in Figure 3-15.

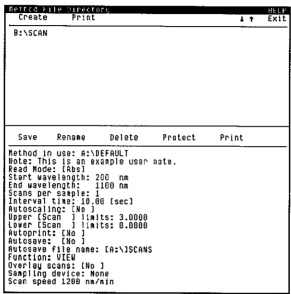


Figure 3-15. Typical Method Window

Each Method window is divided into two parts. The top part is the Method File Directory; the bottom part lists the analysis parameters for the "Method in use".

Creating a Method

To create a new method:

- 1. With the Main window displayed, click on the desired analysis mode. The first window for the analysis mode is displayed.
- 2. Click on <Method> to display the Method window for the analysis mode. The analysis parameters are listed in the lower part of the window.

- 3. If the method to be created is similar to an existing method, click on the existing method name in the directory at the top of the Method window. The existing parameters are displayed on the lower part of the window.
- 4. Click on <Create> to display the Create window, Figure 3-16.

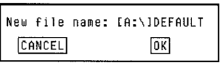


Figure 3-16. Create Window

Input the new method file name and location in this window.

- a. To input the location, click on the displayed location to toggle between [A:\] and [B:\].
- b. To input the new file name, click on the name that is displayed following "New file name" to display the alphanumeric keypad. Input the new name, then click on [OK] to remove the keypad.
- c. Click on [OK] to remove the Create window and accept the new file name.
- d. The new method file name is displayed following "Method in use".
- 5. To input comments about the method, click on "Note" and use the alphanumeric keypad to input a message with a maximum of 40 characters.
- 6. To input the analysis parameters, click on the displayed value(s) and input the desired value(s). Detailed instructions for parameter input are provided in section 3.7.
- 7. After the desired parameters are displayed on the Method window, the method can be used or stored, and if stored, protected.

To use or to store the method, click on < Exit>. The method is stored automatically and the appropriate analysis window is displayed with the parameters for the method.

To store the method without removing the Method window, click on <Save>. This allows more than one method to be created without leaving the Method window.

Protected Methods

After a method has been developed and stored, it can be protected. The parameters for a method that has been protected cannot be changed, as long as the protection remains.

To protect a method:

- Display the desired method name on the Method window, following "Method in use".
- 2. Click on <Protect> to display the Set Protection window, Figure 3-17.

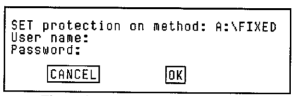


Figure 3-17. Set Protection Window

3. Input the assigned "User name" and "Password", then click on [OK]. If both are accepted, the protection is implemented and the window is removed from the display. **PROTECTED** is displayed following the method name on the Method window.

To remove protection:

Protection is removed using the same steps. If a protected method is displayed following "Method in use", when <Protect> is clicked on, the Remove Protection window shown in Figure 3-18, is displayed. To delete the protection, input the assigned "User name" and "Password", and click on [OK].

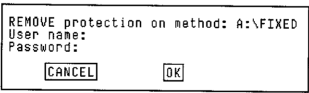


Figure 3-18. Remove Protection Window

Placing the Method Name on the Main Window

The method can be listed on the Main window in the area entitled "Custom Applications". To place it there, use the File Utilities mode to copy the method file from the appropriate method directory to the "CUST_APP" directory. Up to 30 methods can be listed on the Main window.

Using a Stored Method

To use a stored method listed in the "Custom Applications" area on the Main window, click on the desired method name. The appropriate analysis window with the values for the method is displayed.

NOTICE

When a method listed in the "Custom Applications" area is selected, the parameters can be modified on the applications window or the Method window. However, the changes cannot be stored. To store the changes, recall the method from the applications mode, make the changes, store the method, then move it to the Custom Applications directory.

To use a stored method not listed on the Main window:

- 1. With the Main window displayed, click on the desired analysis mode. The first window for the analysis mode is displayed.
- Click on <Method> to display the Method window for the analysis mode.
- 3. The Method File Directory for the analysis mode is displayed at the top of the Method window. Click on the desired method name. The parameters are displayed on the lower part of the window.
- 4. If the method is protected, the protection is indicated to the right of the method name on the lower part of the window. If the method is not protected, any of the parameters can be modified.
- 5. Click on **Exit** to display the appropriate analysis window with the values for the method.

Renaming a Method File

- 1. Display the desired method name, following "Method in use", on the Method window.
- 2. Click on <Rename> to display the Rename window, Figure 3-19.



Figure 3-19. Rename Window

- 3. Click on the name following "New file name" and input the desired file name. Then click on [OK] to rename the file and remove the window from the display.
- 4. The file is stored with the new name when either <Save> or <Exit> (to remove the Method window) is clicked on.

Deleting a Method File

- 1. Display the desired method name, following "Method in use", on the Method window.
- 2. Click on < Delete > to display the Delete window, Figure 3-20.

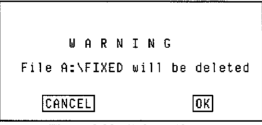


Figure 3-20. Delete Window

3. Verify that the desired file name is displayed, then click on [OK] to delete the file and remove the window from the display.

3.9 Stored Data

The instrument has the capability of storing the data it collects either in internal memory (drive A) or on the optional external disk drive (drive B). Data from most modes are stored in "Results Files". However, some modes have special types of data files, such as "Standard Files" in the protein analysis mode. All types of data files operate the same.

Data files are renamed, copied to another location, moved to another location and deleted using the File Utilities mode. Instructions are provided in section 8.

The following options are available for data storage:

1. Not storing the data.

The data are removed from the window by clicking on either <SaveClear> or <Quit> to display the appropriate window. The data are deleted if the box next to the file name is *not* darkened. The Save Clear window is shown in Figure 3-10. The Quit window is shown in Figure 3-11.

- Designating a file name for data storage before the data are collected.
 The file name is input by clicking on "Results file", displayed with the other parameters on the analysis window.
- 3. Collect the data and then decide whether to store the data.
 When the data are removed from the window by clicking on either <SaveClear> or <Quit>, the appropriate window is displayed.
 The data are stored if the box next to the file name is darkened.

Naming a File

When "Results file" is clicked on from an analysis window, or when the file name is clicked on from the Save Clear or Quit window, the Results File Directory window specific for the analysis mode is displayed. A typical Results File Directory window is shown in Figure 3-21.

To name a new results file:

- 1. To input the location, click on the location that is displayed following "Selected file" to toggle between [A:\] and [B:\].
- 2. To input the new file name, click on the name that is displayed following "Selected file" to display the alphanumeric keypad.

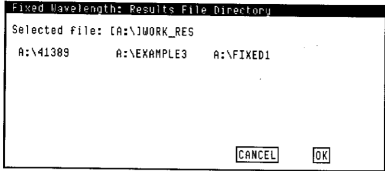


Figure 3-21. Results File Directory Window

- 3. Input the new file name, then click on [OK] to remove the keypad.
- 4. Click on [OK] to remove the window and accept the new file name.
- 5. The new file name is displayed following "Results file".

Recalling Data

Data files that are stored in the instrument can be recalled in the mode where they were created. The stored information can be used to recalculate results in the same way that the data was manipulated after it was collected. With the exception of the Fraction Read mode, data cannot be added to an existing file.

To recall data:

- 1. With the Main window displayed, click on the desired analysis mode. The first window for the analysis mode is displayed.
- 2. Verify that no data are displayed on the window. If data are displayed, click on <SaveClear>.
- 3. Click on the file name following "Results file". The Results File Directory window is displayed.
- 4. Click on the desired file name, listed in the directory to place the name after "Results file name". Click on [OK] to remove the Results File Directory window and display the data in the appropriate analysis window.

GETTING STARTED

The DU Series 600 Spectrophotometer user interface operates on the principle of windows. The "mouse" is used to position an arrow on the window. When the arrow points to the desired position, the left button on the mouse is pressed to initiate the desired action. In these instructions, the positioning of the arrow and pressing the left mouse button is called "clicking on".

4.1 Power Up

Power Up Diagnostics Window

When the DU Series 600 Spectrophotometer is powered up, the Power Up Diagnostics window, Figure 4-1, is displayed. If all tests passed, use the mouse to move the arrow so that it points to "Quit", located near the top right-hand corner of the window, and press the left mouse button.

Power Up Diagnostics Print			Quit
			quic
Computer and Hardware	Diagnost	ics:	
CPU PROM RAM Controller RAM Video Controller Video RAM Video Palette RS232 Ports 1 and EE PROM		Passed Passed Passed Passed Passed Passed Passed Passed Passed	
Spectrophotometer and	Systems	Diagnostic	s:
PROM Option Software Option RAM Option RAM Battery Backup Programmability RS232 Ports 3 and Keyboard Processor Detector Gain Visible Lamp Light Path Filter Lamp Selector Wavelength Drive System Clock	4	Passed Passed Installed Passed	

Figure 4-1. Power Up Window

Getting Started 4-1

NOTICE

If any of these tests fail, refer to the Troubleshooting instructions in section 11.1 of this manual.

Programmability is an option and may not be installed on all instruments.

The "Quit" command is located in the menu bar at the top of the Power Up Diagnostics window. Most windows have a menu bar associated with them. Commands in the menu bar at the top of a window are referred to in these instructions with single angle brackets, i.e. <Quit>.

Main Window

When the Power Up Diagnostics Window is removed, the Main window, Figure 4-2, is displayed. The desired operating mode is selected from the Main window. The Fixed Wavelength, Wavelength Scan and Kinetics/Time modes are standard on all instruments. All other modes are optional, and are only displayed if they are installed on the instrument.

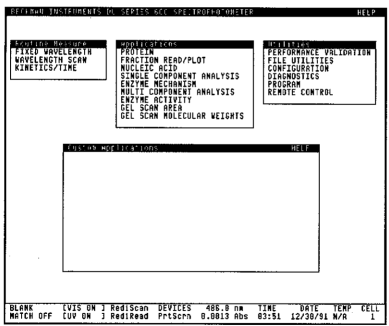


Figure 4-2. Main Window

Sources

The commands to turn the sources on and off are located in the Permanent Menu bar, which is always located at the bottom of the display. Commands in the permanent menu bar are referred to in the instructions with double angle brackets, i.e. <<VIS OFF>>.

To turn on the visible source, click on <<VIS OFF>> to display <<VIS ON>>. The visible source lights immediately.

To turn on the UV source, click on <<UV OFF>> to display <<UV WAIT>>. The UV source requires about 30 seconds to warm up before it lights. When the source lights the command is changed to <<UV ON>>. Do not blank while <<UV WAIT>> is displayed.

The instrument should be allowed to warm up for at least 30 minutes before blanking and taking sample readings. Any reading taken with the sources turned off is invalid.

To turn off the sources, click on <<VIS ON>> and/or <<UV ON>>. When the source is turned back on, a new blank will be required.

Data Collection Modes

The DU Series 600 Spectrophotometer has five data collection modes: RediReadTM Mode, RediScanTM Mode, Fixed Wavelength, Wavelength Scan and Kinetics/Time. They are described in the following sections.

Getting Started 4-3

4.2 RediReadTM Mode

The RediRead window is used to take fixed wavelengths readings at one or more wavelengths quickly and easily. This window can be displayed whenever the instrument is not collecting data, regardless of the operating mode of the instrument. Data collected in this mode cannot be stored.

1. Click on <<RediRead>>, located in the permanent menu bar at the bottom of the display, to display the RediRead window, Figure 4-3.

RediRead ReadSample	ReadBlank	: Print	HELP Exit
500.0nm 0.1	278 A		
Read avg ti Sample Wav		Read Mode: Reading	[Abs]
2 500 3 500		0.0109 A 0.0481 A 0.1091 A 0.1278 A	

Figure 4-3. RediRead Window

- 2. Set the parameters:
 - a. Click on the wavelength value displayed and input the desired wavelength.
 - b. Click on "Read avg time" and input the desired read average time.
 - c. Verify that the desired reading mode is displayed, [Abs] or [%T]. Click on the mode to change it.
- 3. Place a cuvette of solvent in the cell holder and click on <ReadBlank>. (If the instrument has previously been blanked at the selected wavelength using <<BLANK>>, it is not necessary to blank in the RediRead mode. <ReadBlank> in the RediRead mode does not affect the blank stored using <<BLANK>>.)
- 4. Place a cuvette of sample solution in the cell holder and click on <ReadSample>. The reading is displayed in the table on the window.

- 5. Repeat step 4 for all the samples. The parameters input in step 2 can be changed at any time.
 - Readings from 11 samples are displayed on the window. When the sample 12 is read, the data is written over the data for sample 1.
- 6. To print the window, click on <**Print>**. Only the data that are displayed are printed.
- 7. To remove the RediRead window, click on <Exit>.

4.3 RediScanTM Mode

The RediScan window is used to make a wavelength scan at 1200 nm/min on a sample with minimum parameter setup. Data collected using this window cannot be stored; the Wavelength Scan mode must be used for data storage.

1. Click on <<RediScan>>, located in the permanent menu bar at the bottom of the display, to display the RediScan window, Figure 4-4.

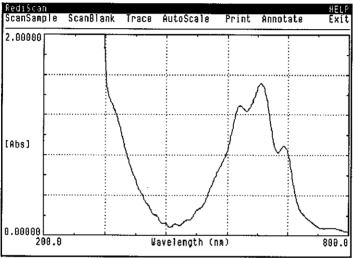


Figure 4-4. RediScan Window

- 2. Verify that the proper ordinate label is displayed, [Abs] or [%T]. Click on the label to change it.
- 3. Verify that the desired wavelength limits are displayed. To change them, click on the displayed value and input the desired value. The sample will be scanned over the displayed wavelength range, only.
- 4. Place a cuvette of solvent in the cell holder and click on <ScanBlank>. (If the instrument has previously been blanked in the Wavelength Scan mode at 1200 nm/min over the selected range, it is not necessary to blank in the RediScan mode.)
- 5. Place a cuvette of sample solution in the cell holder and click on <ScanSample>. The scan data is displayed.

- 6. The following functions are available to reformat the data:
 - a. The data can be autoscaled by clicking on <AutoScale>.
 - b. Individual axis limit values can be changed by clicking on them and inputting the desired value.
- 7. To display the wavelength and ordinate readings at any point in the spectrum, click on <Trace>. Then move the mouse to the point of interest in the spectrum and click on the center mouse button to place a vertical line on the spectrum. The values at the place where the vertical line is placed are displayed in the lower right-hand side of the window. To move the vertical line to either the right or left, click on the right or left mouse button, respectively.
- 8. To annotate the data, click on <Annotate>. Then, click on the graph to position a cross and input information from the alphanumeric keypad or keyboard. Up to four annotations can be placed on the graph. The annotations are printed with the window, but are not stored with the data.
- 9. To print the wavelength scan in the window, click on <Print>.
- 10. Repeat steps 5 to 8 for all the samples.
- 11. To remove the RediScan window, click on **Exit>**.

4.4 Fixed Wavelength

The Fixed Wavelength mode is used to collect data from a series of samples at up to 12 wavelengths. The data can be multiplied by user-input factor(s) to calculate a result at each wavelength. Any of the sampling devices can be used to simplify sample handling. Data can be stored for later recall.

To select the analysis parameters:

1. With the Main window displayed, click on "FIXED WAVELENGTH" to display the Fixed Wavelength window, Figure 4-5.

Fixed Wavelergt ReadSamples	h Method	Parameters	SaveClear	Print	HEL≅ Quit
Results file: Read average t		ead mode: [Abs]	Method name: A:\F Sampling device:		• † •
Sample ID		0.0 λ 56.00 Factor	140.0 λ 230.0 Factor	520.0 6.500	
1 2 43F 43T 46J 48K 7	0.2790 15 0.3647 20 0.6747 37 0.6413 35 1.0447 58	.4213 0.0971 .7840 0.2244 .9108 0.2421 .5056 0.3162	Result Rbs ng/nl 15.2955 8.3152 22.3246 0.3784 11.6812 1.0832 15.6869 6.8864 22.7205 1.4383 16.6482 1.3800	Result Rg/nl 2.4596 2.4596 7.6497 4.4613 9.2369 8.9698	

Figure 4-5. Fixed Wavelength Window

2. Click on <Parameters> to display the Parameters window, Figure 4-6.

Fixed Wavelend ClearAll	ith: Para Print	meters	Exit
Wavelength	Factor	Units	Use
350.0 440.0 520.0 200.0 400.0 250.0 300.0 550.0 600.0 750.0	56.30 230.0 6.500 1.000 1.000 1.000 1.000 1.000 1.000 1.000	mg/ml mg/ml mg/ml mg/ml mg/ml mg/ml mg/ml mg/ml mg/ml	Cyes] Cyes] Cyes] Cho]

Figure 4-6. Parameters Window

- a. Listed in the Parameters window are 12 wavelength values, with a factor and units that correspond to each wavelength. To change any of these values, click on the displayed value to display a keypad. Input the desired value on the keypad, then click on [OK] to accept the input and remove the keypad.
- b. The fourth column in the Parameters window is the "Use" column. Each wavelength that is to be used in the analysis must have a "Yes" displayed. If a "No" is displayed for a desired wavelength, click on the "No" to display a "Yes".
- c. When all the desired values are displayed, click on <Exit> to remove the Parameters window. The input values are immediately displayed on the Fixed Wavelength window.
- 3. Readings can be taken in either absorbance or transmittance. The selection is displayed following "Read mode" in the parameter listing near the top of the window. To change the read mode, click on the displayed option.

To take readings:

- 1. Place a cuvette of solvent in the instrument. Click on <<BLANK>>.
- 2. If desired, click on the next displayed sample number and input up to an 11-digit alphanumeric sample identification. If a sample identification is not input, the instrument numbers the samples consecutively.

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- 3. Place a cuvette of sample solution in the cell holder and click on <ReadSamples>.
- 4. Data from up to 3 wavelengths are displayed at one time. To display data at other selected wavelengths, click on the right and left arrows, located on the right-hand side of the analysis parameters.
- 5. Repeat steps 2 to 4 until all samples have been read.
- 6. To print the sample data, click on <Print>.
- 7. When the analysis is complete, click on **Quit>**. To store the method and/or results, click on the displayed file name(s) and input the desired file name(s). Then click on **[OK]** to store the data and return to the Main window.

The complete capabilities of the Fixed Wavelength mode are described in section 5 of this manual.

4.5 Wavelength Scan

The Wavelength Scan mode is used to collect, manipulate and store scan data.

To select the analysis parameters:

 With the Main window displayed, click on "WAVELENGTH SCAN" to display the Wavelength Scan window, Figure 4-7.

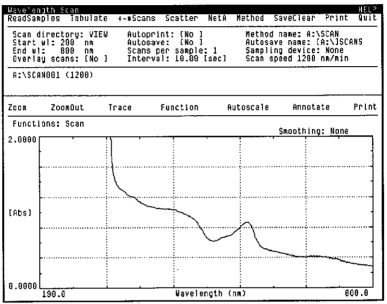


Figure 4-7. Wavelength Scan Window

- 2. Twelve parameters are listed near the top of the window.
 - a. Locate the "Start wl" and "End wl" parameters. To change the values, click on the displayed value and input the desired values.
 - b. Verify that the following parameters are as follows:

Overlay scans: [No]
Autoprint: [No]
Autosave: [No]
Scans per sample: 1
Sampling device: None
Scan speed: 1200 nm/min

If any of the parameters are different than those listed, click on the displayed value and input the listed value.

3. The ordinate label and limits are displayed on the graphic portion of the window. To change any of these values, click on the displayed value and input the desired value.

To take readings:

- 1. Place a cuvette of solvent in the cell holder. Click on <<BLANK>>.
- 2. Place a cuvette of sample solution in the cell holder and click on <ReadSamples>.
- 3. The following functions are available to reformat the data:

Autoscale - Automatically scales the ordinate axis.

Limit changes - The limits on either axis can be changed by clicking on the displayed value and inputting the desired value.

Zoom - The "zoom" feature is used to expand any portion of the graph. Click on <Zoom>, then click on two points on the graph to place crosses at the opposite corners of the area to be enlarged. When the second cross is clicked on, the graph is replotted. This can be repeated as often as desired. To return to the original plot, click on <ZoomOut>.

- 4. To smooth the data, click on "Smoothing" and select the desired number of points to use for the calculation. If too few points are used, the data may appear to be noisy. If too many points are used, real peaks can be lost.
- 5. To display a derivative or log scan, peak pick, valley pick and/or point pick, click on <Function> to display the Function Selection window. To select a function, click on to darken the box preceding the selection. Then click on <Exit> to remove the Function selection window from the display. The data are replotted using the selected function(s).
- 6. To display the wavelength and ordinate readings at any point in the spectrum, click on <Trace>. Then move the mouse to the point of interest in the spectrum and click on the center mouse button to place a vertical line on the spectrum. The values at the place where the vertical line is placed are displayed in the lower right-hand side of the window. To move the vertical line to either the right or left, click on the right or left mouse button, respectively.

- 7. To annotate the data, click on <Annotate>. Then, click on the graph to position a cross and input information from the alphanumeric keypad or keyboard. The annotation is printed with the window, but is not stored with the data.
- 8. To print the sample data, click on < Print>.
- To store data before scanning another sample, click on <SaveClear>.
 Click on the displayed file name, input the desired file name, then click on [OK]. The data are stored and the graphic area is cleared.
- 10. To scan additional samples, repeat steps 2 to 9, above.
- 11. When all the samples have been scanned, click on <Quit>. To store the method and/or displayed scan, click on the displayed file name(s) and input the desired file name(s). Then click on [OK] to store the data and return to the Main window.

The complete capabilities of the Wavelength Scan mode are described in section 6 of this manual. These include selection of different analysis parameters, overlaid scans, repetitive scanning, scatter correction, spectral manipulation (addition, subtraction and multiplication), and net absorbance.

4.6 Time Drive

The Kinetics/Time mode is used to collect, manipulate and store time drive data. This mode is also used to calculate the rate of kinetic reactions.

To select the analysis parameters for time drive:

1. With the Main window displayed, click on "KINETICS/TIME" to display the Plotting window, Figure 4-8.

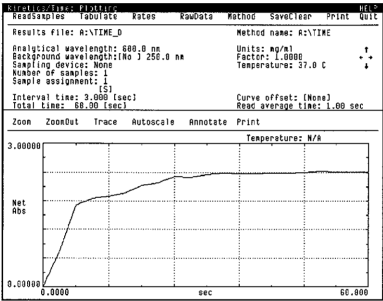


Figure 4-8. Plotting Window

- 2. Fourteen parameters are listed near the top of the window.
 - a. Locate "Analytical wavelength". To change the wavelength, click on the displayed value and input the desired wavelength.
 - b. Locate "Interval time", which determines the frequency of data collection. To change the displayed value, click on it and input the desired value.
 - c. Locate "Total time", which determines when data collection is stopped. To change the displayed value, click on it and input the desired value.

d. Verify that the following parameters are as follows:

Background wavelength: [No]

Sampling device: None Number of samples: 1 Sample assignment: [S] Read average time: 0.5 sec

If any of the parameters are different than those listed, click on the displayed value and input the listed value.

3. The absorbance limits are displayed on the graphic portion of the window. To change the limits, click on the displayed value and input the desired value.

To take readings:

- 1. Place a cuvette of solvent in the cell holder. Click on <<BLANK>>.
- 2. Place the sample in the cell holder and click on <ReadSamples>. The Read Samples window is displayed. Click on [START].
- 3. The data are displayed as they are collected. If no data appear on the graph, the data probably do not fall within the axis limits.
- 4. After data collection is complete, the following functions are available to reformat the data:

Autoscale - Automatically scales the ordinate axis.

Limit changes - The limits on either axis can be changed by clicking on the displayed value and inputting the desired value.

Zoom - The "zoom" feature is used to expand any portion of the graph. Click on <Zoom>, then click on two points on the graph to place crosses at the opposite corners of the area to be enlarged. When the second cross is clicked on, the graph is replotted. This can be repeated as often as desired. To return to the original plot, click on <ZoomOut>.

- 5. To annotate the data, click on <Annotate>. Then, click on the graph to position a cross and input information from the alphanumeric keypad or keyboard. The annotation is printed with the window, but is not stored with the data.
- 6. To print the sample data, click on **<Print>**.

Getting Started 4-15

- 7. To display the tabulated data, click on <Tabulate>.
- 8. Before analyzing another sample, click on **SaveClear**. To store the data, if desired, click on the displayed file name and input the desired file name, then click on [OK]. The data are stored and the graphic area is cleared.
- 9. To analyze additional samples, repeat steps 2 to 8, above.
- 10. When all the samples have been analyzed, click on **<Quit>**. To store the method and/or displayed data, click on the displayed file name(s) and input the desired file name(s). Then click on **[OK]** to store the data and return to the Main window.

The complete capabilities of the Kinetics/Time mode are described in section 7 of this manual. These include selection of different analysis parameters, analysis of multiple samples, and rate calculations.

4.7 Recalling Stored Files

Method File

If a method was stored in one of the operating modes, it can be recalled by clicking on the file name following "Method name", listed with the analysis parameters at the top of the analysis window. When the file name is clicked on, the Method window is displayed. The stored method files are displayed near the top of the window. To select a stored file, click on the file name. The parameters will be listed in the lower portion of the window. Then click on <Exit> to remove the method window and display the analysis parameters on the analysis window.

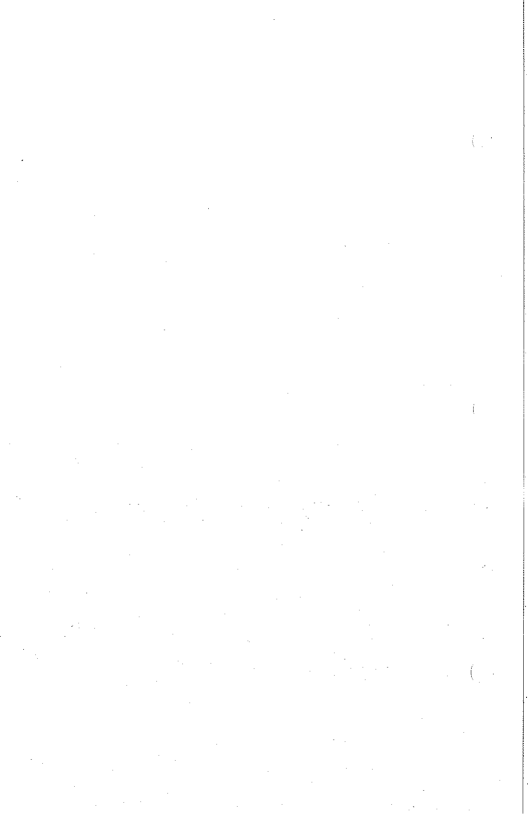
Results File - Fixed Wavelength and Kinetics/Time

If a results file was stored in the Fixed Wavelength or Kinetics/Time mode, it can be recalled by clicking on the file name following "Results file" on the analysis window. When the file name is clicked on, the directory is displayed. To select a stored file, click on the file name, then click on [OK] to remove the directory and display the data on the analysis window.

Scan File - Wavelength Scan

If a scan file was stored, it can be recalled by clicking on "VIEW", following "Scan directory" on the Wavelength Scan window. When the file name is clicked on, the directory is displayed. To select a stored file, click on the file name, then click on [OK] to remove the directory and display the data on the Wavelength Scan window.

Getting Started 4-17



FIXED WAVELENGTH

The Fixed Wavelength mode is used to collect data in either absorbance or transmittance at up to 12 wavelengths. The reading at each wavelength can be multiplied by a user input factor to calculate a final result.

5.1 Calculations

The result is calculated using the equation:

Result = Reading x Factor,

where the reading is in either absorbance or transmittance. The result is a concentration value if the reading is taken in absorbance.

NOTICE

Use of this mode to calculate concentration requires that the slope of the standard curve is constant and known, and that the y-intercept is zero. Concentration calculations, derived from a standard curve with multiple standards, are possible using the optional Single Component Analysis mode.

5.2 Parameter Setup

Click on "FIXED WAVELENGTH" from the Main window to start the analysis. The Fixed Wavelength window, Figure 5-1, is displayed. The Fixed Wavelength window is used to select analysis parameters, collect sample data and display stored sample data.

n Method	Paras	neters	Sa	vellear	Print	4ELP Quit
a:\WORK_RES iae: 0.50	Read Rode	e: [Abs]	Sampling	davice:	EFAULT None	†
λ 20 Factor	1.000	λ 2 Factor	50.0 1.000	λ 3 Factor	1.000	
Abs	Result mg/ml	Abs	Result mg/ml	Abs	Result mg/ml	
	Method A:\⊎ORK_RE iae: 0.50 A 20 Factor	Method Para A:\WORK_RES ine: 0.50 Read Rodo \(\lambda\) 200.0 Factor 1.000 Abs Result	Method Parameters A:\Work_RES ime: 0.50 Read mode: (Abs) \[\frac{206.0}{Factor} \frac{22}{1.000} 22	Method Parameters Sa a:\Work_RES Method no ime: 0.50 Read mode: (Abs) Sampling Factor: \$\text{\lambda} 200.0 \text{\lambda} 250.0 \$\text{\lambda} Factor 1.000 Factor 1.000 \$\text{\lambda} Result Abs Result	Method Parameters Saveflear A:\WORK_RES	Method Parameters Saveflear Print A:\WORK_RES

Figure 5-1. Fixed Wavelength Window

Use the Method window to setup the analysis parameters:

- Click on <Method> to display the Method window, Figure 5-2. The Method window is used to setup analysis parameters, recall stored methods and create new methods. General information on method windows is provided in section 3.8.
- 2. To recall a stored method, click on the desired method name in the listing at the top of the Method window.
- 3. The analysis parameters are displayed on the lower part of the Method window. Input the desired analysis parameters:

Method in use - This displays the name of the method that has been selected. If the method is protected, **PROTECTED** is displayed following the method name. If the method is protected, the analysis parameters cannot be changed. To input a new method name, click on <Create>.

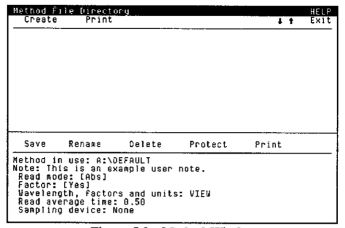


Figure 5-2. Method Window

Note - Click on to input a 40-character message that is used to describe the method or procedure.

Read mode - Toggle between [Abs] and [%T] to select readings in absorbance or transmittance, respectively.

Factor - Toggle betweem [Yes] and [No] to specify whether a factor and units are input for each wavelength for results calculations.

Wavelength, factors and units - Click on "VIEW" to display the Parameters window, Figure 5-3. The Parameters window is used to input analysis wavelength(s), a concentration factor for each wavelength and the concentration units for each wavelength.

Fixed Wavelend ClearAll	ith: Para Print	meters	Exit
Wavelength	Factor	Units	Use
200.0 250.0 350.0 400.0 450.0 550.0 650.0 700.0	1.000 1.000 1.000 1.000 1.000 1.000 1.000 1.000 1.000 1.000	mg/ml mgg/ml mgg/ml mgg/ml mg/ml mg/ml mg/ml mg/ml	Yes] Yes] Yes] Yes] You I Yes] Yes] Yes] Yes] Yes] Yes] Yes] Yes]

Figure 5-3. Parameters Window

Fixed Wavelength 5-3

- a. Input the desired values for wavelengths, factors and units by clicking on the respective location and inputting the desired values. Up to 12 wavelengths can be input.
- b. Enable or disable each wavelength by clicking on [Yes]/[No] in the "Use" column.
- c. When all the desired values are input, click on **Exit** to accept the values and remove the window from the display.

Sampling device - Display the Sampling Device window to select the sampling device. Detailed information on the sampling devices is provided in Manual 517314.

Read average time - The time, in seconds, that data are collected and averaged to take a reading. Twenty sets of data are collected every second.

- 3. To store the analysis parameters in the selected method file, click on <Save>.
- 4. Click on **Exit** to display the Fixed Wavelength window with the parameters from the selected method.

To input the desired parameters on the Fixed Wavelength window:

- 1. The analysis parameters are listed near the top of the window. To change any of these, click on the parameter and input the desired value. A description of the parameters is provided above.
- 2. To change the number of wavelengths, the wavelength value, factor or concentration units, click on <Parameters> to display the parameters window, Figure 5-3. As an alternative, the wavelength, factor and concentration units, displayed in the table on the Fixed Wavelength window, can be changed by clicking on them and inputting the desired values. The Parameters window must be used to change the total number of enabled wavelengths.

5.3 Sample Analysis

After the desired parameters are displayed on the Fixed Wavelength window, samples can be run.

1. Place a cuvette of solvent in the cell holder. Click on <<BLANK>>. The instrument blanks on the solvent.

<u>Auto Cell Holder</u> - Place the cuvette of solvent in the cell position that is in the light beam. Click on **<<BLANK>>**.

<u>Sipper/Batch Sampler</u> - Aspirate and read the solvent by pressing {FILL/BLANK}.

- If desired, click on the file name following "Results file" and input the desired file name. As an alternative, the file can be named after the data are collected.
- 3. If desired, click on the next displayed sample number to display the alphanumeric keypad. Input up to an 11-character alphanumeric sample identification using the mouse, the optional keyboard or the optional Bar Code Accessory. If a sample identification is not input, the instrument numbers the samples consecutively.

NOTICE

The sample number can be input only for the next sample to be analyzed. These values cannot be changed for samples that have already been analyzed.

4. Place a cuvette of sample solution in the cell holder. Click on <Read-Samples>. (As an alternative, click on the right mouse button with the cursor in any position.)

<u>Auto Cell Holder</u> - Place the cuvettes of sample solution into the input number of cell positions in the sampler. Click on **<ReadSamples>**.

<u>Sipper</u> - Aspirate and read each sample solution by pressing {FILL/READ}. Flush or return each sample solution before reading the next.

ISCO Sampler - Load the tubes of sample solutions into the batch sampler, starting with the position directly under the aspirator arm. Click on <ReadSamples>. The sample solutions are read until the last position of the red rack is read.

<u>Gilson Sampler</u> - Program the Sample Controller with the rack type, number of tubes and orientation of the tubes. Load the samples in the programmed configuration. Click on <ReadSamples>. The analysis stops when the programmed number of tubes are read.

- 5. Data from up to 3 wavelengths are displayed at one time. To display data at other selected wavelengths, click on the right and left arrows, located on the right-hand side of the analysis parameters.
- 6. Repeat steps 3 to 5 until all samples have been read. A typical Fixed Wavelength window is shown in Figure 5-4.

ixed Naveleno eadSamples	th Method	Par	ameters	S	aveClear	Print	HEL Quí
Results file: Read average		Read no	de: [Abs]		name: A:\ g device:		-1
Sample ID	Factor	350.0 56.08	λ Facto	440.0 r 230.0	λ Facto	520.0 r 6.500	
	Abs	Result	Abs	Result ng/ml	Abs	Result mg/ml	
1 2	0.3647	15.6221 20.4213	0.1535 0.0971	35.2955 22.3246	0.3152 0.3784	2.0499 2.4596	
43F 43T	0.6413	37.7848 35.9108	0.2244 0.2421	51.6012 55.6869	1.0832 0.6864	7.0487 4.4613	
46J 48K 7		58.5056 53.2240	0.3162 0.3767	72.7205 86.6482	1.4303 1.3800	9.2969 8.9698	
ſ							

Figure 5-4. Fixed Wavelength Window

- 7. To display sample data that has scrolled off the window, click on the arrows, located on the right-hand side of the analysis parameters, to scroll through the data.
- 8. To print the sample data, click on < Print>. All sample data, even that which are not displayed because of insufficient room on the display, are printed.

NOTICE

When analyzing at more than three wavelengths, the condensed print mode on the printer is suggested for the optimum data formatting. 9. To clear all sample data from the window or to store the data, then clear the window, click on **SaveClear>**. The Save Clear window is displayed so the sample data can be stored.

NOTICE

Do not store the data until all sample data have been collected. After the data are stored, no additional data can be placed in the Results file.

10. When the analysis is complete, click on <Quit>. The Quit window is displayed so the method and/or sample data can be stored. Then the Main window is displayed.

5.4 Example Analyses

EXAMPLE 1

Create a method to collect data on chromate samples at 500 nm with a read average time of 1.0 second. Multiply the readings by a factor of 125 to calculate the concentration in units of $\mu g/mL$.

SOLUTION

The Method window shown in Figure 5-5 was used to create this method. The name of the method is CHROMATE and it is stored on the A drive.

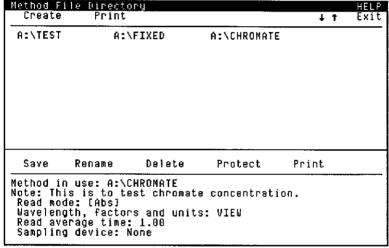


Figure 5-5. Method Window for Chromate Analysis

EXAMPLE 2

Run the method developed in EXAMPLE 1. Analyze four samples, with sample numbers of 41389.1, 41389.2, 41389.3 and 41389.4. Store the data in a file on the optional B drive named 41389.

SOLUTION

The method named "A:\CHROMATE" was recalled into the Fixed Wavelength window, shown in Figure 5-6. The results file was named "B:\41389". Sample readings were taken by clicking on <ReadSamples>.

med Wavelen adSamples	Method	Parameters	SaveClear	Print	Gui
lesults file: lead average	B:\41389 time: 1.00	Read mode: [Abs]	Method name: A:\CHR Sampling device: No		† + 1
Sample IĐ	Factor	500.0 125.0			
	Abs	Result			
41389.1	0.1835	ug/mL 22 9426			
41389.2	0.2176	22.9426 27.2048			
41389.3	0.2722	34,0193			
41389.4	0.2132	26.6459			
5					

Figure 5-6.
Sample Data for Analysis of CHROMATE Method

EXAMPLE 3

Using the Fixed Wavelength window, setup and run an analysis of samples at the following wavelengths: 230, 260, 280 and 320 nm. Use the sipper for the analysis. Do not store the data.

SOLUTION

The parameters were input into the Parameter window by clicking on <Parameters> from the Fixed Wavelength window. The results file was named "WORK_RES" because the data were not stored. The data are shown in Figure 5-7. Part of the data at 320 nm is not displayed. Click on the right arrow to display the remainder of the data.

Fixed Waveleng ReadSamples	th Method	Par	ameters	S	aveClear	Pri	HELP nt Quit
Results file: Read average			de: [Abs]		name: A:\ g device:		↑
Sample ID	Factor	230.0	λ Factor	268.0 1.000	Facto	280.0 r 1.000	λ Fact
	Abs	Result ลญ/ตไ	Abs	Result ng/ml	Abs	Result ag/ml	Abs
1 2 3 3 4 4 5 6	0.3015 0.6146 0.8658 0.1983 1.5613	9.3915 9.8146 9.8658 9.1963 1.5613	0.2865 0.5928 0.8924 0.2041 1.6238	9.2965 9.5928 9.8924 9.2941 1.6238	0.2824 0.5692 0.9189 0.2074 1.6190	0.2824 0.5692 0.9109 0.2074 1.6190	8.2711 9.4563 9.7929 9.2847 1.4864

Figure 5-7. Analysis of Samples at 4 Wavelengths

5.5 Data Output

Data are output to the Communications port whenever an Output Data Type is selected in the Communications Configuration window. The type of data output is selected: "raw and calculated", "raw", or "calculated". General information on data output is provided in section 9.3.

In the Fixed Wavelength mode, the order of the output for each data type is shown below. The "User ID" is input in the Communications Configuration window. The "Method ID" is the method name. A <carriage return> and are sent at the end of every line.

Raw and Calculated

The data output for each sample includes the sample number followed by the reading and calculated result for each wavelength. The following is the output obtained when the data in Figure 5-4 were collected.

User ID: [-					
	: A:\FIXED					
Date: 10\(09\91					
Time: 10:1	18					
1	0.2790	15.6221	0.1535	35.2955	0.3152	2.0490
2	0.3647	20.4213	0.0971	22.3246	0.3784	2,4596
43F	0.6747	37.7840	0.2244	51.6012	1.0832	7.0407
43T	0.6413	35.9108	0.2421	55.6869	0.6864	4.4613
46J	1.0447	58.5056	0.3162	72.7205	1.4303	9.2969
48K	0.9584	53,2248	0.3767	86,6482	1.3800	8.9698

Fixed Wavelength 5-11

Raw

The data output for each sample includes sample number followed by the reading at each wavelength. The following is the output obtained when the data in Figure 5-4 were collected.

User ID: D	U600		
Method ID:	A:\FIXED		
Date: 10\0	9\91		
Time: 10:1	8		
1	0.2790	0.1535	0.3152
2	0.3647	0.0971	0.3784
43F	0.6747	0.2244	1.0832
43T	0.6413	0.2421	0.6864
46J	1.0447	0.3162	1.4303
48K	0.9584	0.3767	1.3800

Calculated

The data output for each sample includes the sample number followed by the result at each wavelength. The following is the output obtained when the data in Figure 5-4 were collected.

```
User ID: DU600
Method ID: A:\FIXED
Date: 10\09\91
Time: 10:18
                15.6221
                         35.2955
                                   2.0490
2
                20.4213 22.3246
                                   2,4596
43F
                37.7840 51.6012
                                   7.0407
                35.9108
43T
                         55,6869
                                   4.4613
46J
                58.5056 72.7205
                                   9.2969
                53.2248 86.6482
48K
                                   8.9698
```

5.6 Files

Two file types are created in the Fixed Wavelength mode: method files and data files. The method files are stored in the FIX_METH directory with the .APX extension. Method files are ASCII files and cannot be converted to the Lotus format.

The data files are stored in the FIX_DATA directory with the .DUF extension. They contain analysis parameters and sample data.

5.7 ASCII Format

The ASCII file for the Fixed Wavelength mode consists of two parts: analysis parameters and sample data, in the same format as the Fixed Wavelength window.

The analysis parameter, "Mode", is either 0 (absorbance) or 1 (transmittance).

The sample data include: sample identification and the reading and calculated result at each wavelength.

The following ASCII file was converted from the sample data displayed in Figure 5-4.

Mode:	0					
	r of waveler	naths: 3				
		Factor				
	0.0000 mg/ml					
520	0.0000 mg/ml	6.5000				
1	1.2308	68.9246	0.9970	229.3057	0.6357	4.1318
	4 (5/0	92.7359	0.7138	164.1711	0.5403	3.5118
2	1.6560					
2 43F	1.5952	89.3310	1.2345	283.9270	1.0218	6.6416
					1.0218	6.6416 6.4615
43F	1.5952	89.3310	1.2345	283.9270		

Fixed Wavelength 5-13

5.8 Lotus Format

The Lotus file for the Fixed Wavelength mode consists of two parts: analysis parameters and sample data, in the same format as the Fixed Wavelength window.

The analysis parameter, "Mode", is either 0 (absorbance) or 1 (transmittance).

The sample data start in row 9 and are stored in the following columns:

Column A Sample identification

Column B Reading at the first wavelength Column C Result at the first wavelength

Columns D, E Reading and result at the second wavelength, etc.

The following Lotus file was converted from the sample data displayed in Figure 5-4.

```
2
3
4
5
    Number of wavelengths:
    Wavelength
                               Units
                          350
                                  mg/ml
                                              56
                                             230
                           440
                                  mg/ml
                          520
                                  mg/ml
              0.278965
                       15.62207
                                0.153458 35.29548 0.315226 2.048974
10
11
              0.364666
                       20.42131 0.097063 22.32464 0.378403
    43F
                       37.78400 0.224353 51.60123
                                                   1.083182
                       35.91077 0.242117 55.68693 0.686351
    43T
               641263
    46J
                       58.50556 0.316176 72.72050
                                                   1.430297
                950428 53.22399 0.376731 86.64817 1.379972 8.969822
```

SECTION SIX

WAVELENGTH SCAN

Wavelength scans can be collected in either absorbance or transmittance. The spectra are available for formatting changes which include expansion and contraction of either axis, zoom, overlaying of up to twelve spectra, overlaying of multiple functions of the same spectrum, and "Trace" to find the abscissa and ordinate values at any user-selected position.

Calculations can be made on wavelength scan data to find the peaks, valleys and/or specified points, to calculate first to fourth derivative, and to calculate the log of absorbance. Scatter correction can be performed on any spectrum. Spectral addition, subtraction, and multiplication can also be performed on stored spectra. Net absorbance calculations can be made on any stored spectrum using either one or two baseline points.

6

6.1 Principles of Operation

Blank - Background Data

The background data are collected when the instrument is blanked. A cuvette filled with the solvent used to prepare the samples should be in the sample compartment when the blank is read.

The blank data are stored in the instrument and can be reused for an unlimited number of sample spectra. A new blank should be read every time a solvent is changed. For optimum performance, a new blank should also be read if the instrument has not been used for over an hour.

For very high precision data, the same cuvette should be used for the blank and sample spectra, as even matched cuvettes have a slight mismatch which can be seen by the DU Series 600 Spectrophotometer. The blank should be rerun for every sample.

Sample Data

After the sample data are collected, the blank is subtracted and the difference in absorbance (or ratio in transmittance) is plotted on the display and stored in the instrument memory. The data points are connected.

Repetitive Scanning

Repetitive scans can be made by inputting more than one for the "scans per sample" and setting the "interval time" to the desired value. The interval time is the time between the start of successive scans. The minimum interval time is 0.1 seconds.

Derivative

First to fourth derivative can be calculated and displayed on any absorbance spectra using the Savitzky and Golay¹ coefficients. Two data points before and two data points after the point of interest are used to calculate the derivative. (For the fourth derivative, only, three points before and after the point of interest are used.) This calculation is done for every data point in the spectrum to determine the derivative spectrum.

¹Savitzky, A., and Golay, M., Anal Chem 36, p1627f (1964).

Log of Absorbance

The log base 10 is calculated and displayed for absorbance spectra. Because the log of 0 is negative infinity, if some data points in the spectrum are zero or are very close to zero, the log spectrum may not be usable in these regions.

Peak/Valley Pick

The most intense peaks and/or valleys are found and displayed with the spectral data when the peak pick calculation is selected. These are found by calculating the first derivative as described above. When the sign of the derivative changes from positive to negative, a peak is detected. A valley is detected when the sign of the derivative changes from negative to positive. Up to 30 peaks (valleys) can be input. If the instrument finds more than the input number of peaks (valleys), the most intense ones will be tabulated. Each time that the wavelength limits or the smoothing function is changed, the instrument performs the peak pick calculations again.

Point Pick

Point pick is a tabulation of data at up to 12 user-selected wavelengths.

Trace

The actual stored data points are accessed using "Trace".

Smoothing

The displayed wavelength scan can be smoothed using a selectable smoothing function. The calculation, using the Savitzky and Golay¹ method (as modified for end points by Peter A. Gorry²), is done for every data point in the scan, using the data points before and after the point of interest. The user selects the total number of data points used for the calculation, from 7 to 25. Use of too few points may not reduce the spectral noise to the desired level, while use of too many points can cause real peaks to be flattened and/or combined.

²Gorry, Peter A., Anal Chem, 62, 1990, p570f.

Addition/Subtraction/Multiplication - (+-* Scans)

Stored spectra can be added, subtracted or multiplied, using the equation:

Scan =
$$(\text{scan A x F1}) + /-/* (\text{scan B x F2}) + /-/* F3,$$

where scan A and scan B are stored spectra, F1 and F2 are factors which are multiplied by each data point in scan A and scan B, respectively. F3 is an additional value that is added/subtracted/multiplied to each point in the spectrum.

Scatter Correction (Spectral)

Turbid samples, rough samples or particles in a sample cause light from the source to be scattered, so that it does not reach the detector. In the spectrum of a sample with scattering particles, the scattered light appears as increased absorbance with decreasing wavelength. If two points in the spectrum can be identified, where the observed absorbance is entirely a result of scatter, a spectrum of the scatter can be estimated and subtracted from the sample spectrum to produce a sample spectrum in the absence of scatter.

The equation used to estimate the scatter is:

$$A_a = a \lambda^b$$

where,

 $A_{\rm s}$ is the absorbance due to scatter at a particular wavelength, and a and b are constants.

For the scatter correction to be accurate, it is necessary to correctly specify two wavelengths where the observed absorbance is entirely a result of scatter. These points should be at least 50 to 100 nm apart for the most accurate correction.

The spectrum due to scatter and the corrected sample spectrum are displayed, along with the constants "a" and "b".

Net Absorbance Calculations

It is possible to correct a sample absorbance peak for a raised or sloping baseline, which may be caused by turbidity in the sample, using net absorbance calculations. Net absorbance calculations can also correct for small differences in cuvettes caused by slight uncleanness, as shown in Beckman publication T-1548.

The user can choose either one or two baseline points for the calculation. If one baseline point is chosen, the absorbance at the base wavelength is subtracted from the absorbance at the peak wavelength to provide the net absorbance. The base wavelength can be either to the right or left of the peak wavelength.

If two baseline points are chosen, a straight line is drawn between the absorbance values at two base wavelengths. The absorbance of that line at the peak wavelength is subtracted from the absorbance of the sample at the peak wavelength to provide the net absorbance. The base wavelengths can be on either side of the peak wavelength, or can both be on the same side.

Scan Files

Spectral data are stored in a Scan file. Each Scan file contains the spectral data from one sample. The Scan file is named using up to five characters, allowing the instrument to append the file name with three characters. These characters are numbers, starting with 001. For example, if the Scan file is named "SCANS", the data from the first sample will be stored in file "SCANS001". The data from the second sample will be stored in file "SCANS002". Data from up to 999 samples can be stored with the same file name.

If "Autosave" is enabled, the data for every sample is stored automatically, using the input Scan file name. If "Autosave" is not enabled, the data for each sample is stored in a temporary file named "WORK_00X". When both "Autosave" and "Autoprint" are disabled, scans from up to twelve samples are stored in this temporary manner; then the scans must be cleared or saved before additional scans can be collected.

Wavelength Scan 6-5

6.2 Parameter Setup

Click on "WAVELENGTH SCAN" from the Main window to start the analysis. The Wavelength Scan window, Figure 6-1, is displayed. The Wavelength Scan window is used to select analysis parameters, collect spectral data and display spectral data. Multiple functions of the sample (i.e. absorbance and first derivative) are displayed in this window. Peak pick, valley pick, and point pick values are also displayed.

NOTICE

Use the RediScan mode to collect and display spectral data with minimum parameter setup.

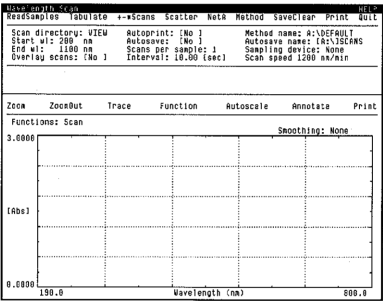


Figure 6-1. Wavelength Scan Window

Use the Method window to setup the analysis parameters:

- 1. Click on <Method> to display the Method window, Figure 6-2. The Method window is used to setup analysis parameters, recall stored methods and create new methods. General information on method windows is provided in section 3.8.
- 2. To recall a stored method, click on the desired method name in the listing at the top of the Method window.

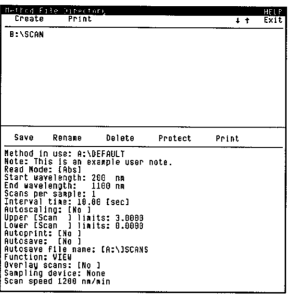


Figure 6-2. Method Window

3. The analysis parameters are displayed on the lower part of the Method window. Input the desired analysis parameters:

Method in use - This displays the name of the method that has been selected. If the method is protected, **PROTECTED** is displayed following the method name. If the method is protected, the analysis parameters cannot be changed. To input a new method name, click on <Create>.

Note - Click on to input a 40-character message.

Read mode - Toggle between [Abs] and [%T] to select readings in absorbance or transmittance, respectively.

Start / End wavelength - Click on to input the starting (lower), then the ending (higher) wavelength.

Scans per sample - For repetitive scanning, input the total number of times each sample is to be scanned. If the Auto Cell Holder or Batch Sampler is selected for the "Sampling device", the input for this parameter is set to 1.

Interval time - For repetitive scanning, input the time between the start of successive scans, in either seconds or minutes. The

Wavelength Scan 6-7

minimum interval time is dependant upon the wavelength range and whether "Autosave" and/or "Autoprint" are enabled.

Autoscaling - Autoscaling adjusts the ordinate axis to display the highest peak full-scale. Toggle between [Yes] and [No] to enable or disable autoscaling.

Upper / Lower limit - Input the upper and lower limits for the ordinate axis display of the spectral data. If autoscale is enabled, the displayed values are ignored.

Autoprint - Toggle between [Yes] and [No]. If enabled, each spectrum is printed automatically on the Dot Matrix Printer after data collection is complete.

Autosave - Toggle between [Yes] and [No]. If it is enabled, each scan is stored automatically as it is collected using the input "Autosave file name".

Autosave file name - If autosave is enabled, input a five-character prefix for the file name.

Function - The Function window is used to select derivative or log spectral display and to select peak, valley and/or point pick tabulation. Click on "VIEW" to display the Function window. Refer to section 6.6 for more information. (On the Wavelength Scan window, click on <Function> in the menu bar to display the Function window.)

Overlay scans - Toggle between [Yes] and [No]. If it is enabled, multiple scan files can be overlaid. If it is disabled, only one scan file can be displayed.

Sampling device - Display the Sampling Device window to select the sampling device.

Scan speed - Select from [2400], [1200], [600], [240] and [120] to set the scan speed in nanometers per minute.

- 3. To store the analysis parameters in the selected method file, click on <Save>.
- 4. Click on **Exit>** to display the Wavelength Scan window with the parameters from the selected method.

To input the desired parameters on the Wavelength Scan window:

- 1. The analysis parameters are listed near the top of the window. To change any of these, click on the parameter and input the desired value. A description of the parameters is provided above.
- 2. The ordinate limits are input by clicking on the displayed values on the graph and inputting the desired values.
- 3. Calculations (derivative, peak pick, point pick, etc.) performed on the data are selected by clicking on **Function** to display the Function window. Refer to section 6.6 for more information.

Wavelength Scan 6-9

6.3 Analysis of Single Samples

After the desired parameters are displayed on the Wavelength Scan window, samples can be run.

1. Place a cuvette of solvent in the cell holder. Click on <<BLANK>>. The instrument blanks on the solvent.

Sipper - Aspirate and read the substrate by pressing {FILL/BLANK}.

NOTICE

If the solvent (blank solution) has significant absorbance in the wavelength region of interest, the dynamic range of the instrument can be increased by adjusting the scan gain for the blank. Instructions are provided in section 2.5.

2. Place a cuvette of sample in the cell holder and click on <Read-Samples>. (As an alternative, click on the right mouse button with the cursor in any position.)

<u>Sipper</u> - Aspirate and read each sample by pressing {FILL/READ}. Flush or return each sample before reading the next.

- 3. The sample is scanned and the data is displayed. The data may be stored or printed automatically, based upon the selections for "Autosave" and "Autoprint". A typical Wavelength Scan window with data from a single sample is shown in Figure 6-3.
- 4. If "Autoprint" is disabled, click on <Print> to print the sample data. The data are plotted on the Dot Matrix Printer unless the X-Y Plotter is installed and selected for "Send graph to" in the Printer and Plotter Configuration window.
- 5. The data can be manipulated using the instructions provided in section 6.5. **Function>** can be used to display additional functions (derivative, peak pick, point pick, etc.) using the instructions provided in section 6.6.
- 6. If "Overlay" is enabled, click on **SaveClear>** to remove the data from previous sample(s) before scanning the next sample, if desired.

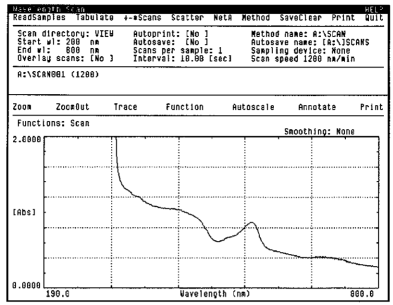


Figure 6-3. Wavelength Scan Window

7. To scan the next sample, repeat steps 2 to 6.

The Save Clear window is displayed after 12 samples are read, unless "Autosave" or "Autoprint" are enabled. The scan data must be stored or cleared before additional samples can be scanned.

8. To exit the Wavelength Scan mode when all spectra have been collected, click on <Quit>. The Quit window is displayed so the method and/or sample data can be stored. Then the Main window is displayed.

Wavelength Scan

6.4 Analysis of Multiple Samples

Wavelength scanning analysis of multiple samples can be automated with the use of a multi-position Auto Cell Holder or one of the batch sampler. When analyzing multiple samples, special considerations should be given to the selection of the optimum analysis parameters.

If an Auto Cell Holder is used, the data can be stored automatically by setting "Autosave" to [Yes] and inputting a file name, or the user has the option of storing the data after they are collected. Regardless of whether they are stored with a permanent file name, they are stored with a temporary identification and can be manipulated after the analysis is complete.

If the batch sampler is used, "Autosave" can also be used to store the data, if sufficient memory exists. However, if "Autosave" is not enabled, the scan data are not stored and therefore the data must either be printed or output to the communications port as they are collected.

To print the data as they are collected, set "Autoprint" to [Yes] and verify that the Dot Matrix Printer is selected for "Send graph to" in the Printer and Plotter Configuration window. To format the data in the desired manner, the following should be considered:

If the data are to be plotted individually, set "Overlay" to [No]. To obtain the optimum plot, set "Autoscaling" to [Yes]. Peak pick can be selected on the Function window and printed with the spectral data.

If the data are to be overlaid, set "Overlay" to [Yes]. "Autoscaling" can also be set to [Yes] to obtain the optimum presentation. Peak, valley, and point pick cannot be selected. Spectra from 12 samples will be overlaid on a single plot, unless less than 12 samples were analyzed.

After the desired parameters are displayed on the Wavelength Scan window, samples can be run.

1. To read the blank:

<u>Auto Cell Holder</u> - Place the a cuvette of blank in the cell position that is in the light beam, then click on **<<BLANK>>**. The instrument blanks on the solvent.

<u>Sipper/Batch Sampler</u> - Aspirate and read the blank by pressing {FILL/BLANK}. The instrument blanks on the solvent.

NOTICE

If the blank solution has significant absorbance in the wavelength region of interest, the dynamic range of the instrument can be increased by setting the instrument gain for the blank. Instructions are provided in section 2.5.

2. To read the samples:

<u>Auto Cell Holder</u> - Place the input number of samples in the sampler and click on <**ReadSamples>**.

<u>ISCO Sampler</u> - Load the samples into the batch sampler, starting with the tube directly under the aspirator arm. Click on **<ReadSamples>**. The samples are read until the last position of the red rack is read.

Gilson Sampler - Program the Sample Controller with the rack type, number of tubes and orientation of the tubes. Load the samples in the programmed configuration. Click on <ReadSamples>. The analysis stops when the programmed number of tubes are read.

3. The samples are scanned and the data are displayed. The data may be stored or printed automatically, based upon the selections for "Autosave" and "Autoprint". A typical Wavelength Scan window with overlaid data from multiple samples is shown in Figure 6-4.

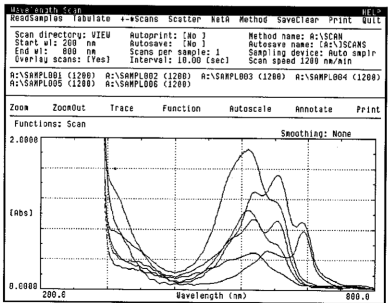


Figure 6-4. Wavelength Scan Window

- 4. If the data are stored, they can be manipulated using the instructions provided in section 6.5. < Function > can be used to display additional functions (derivative, peak pick, point pick, etc.) using the instructions provided in section 6.6.
- 5. If the Auto Cell Holder is used for the sampling device, and both "Autosave" and "Autoprint" are disabled, the Save Clear window is displayed after 12 samples are read. The scan data must be stored or cleared before additional samples can be scanned.
- To exit the Wavelength Scan mode when all spectra have been collected, click on <Quit>. The Quit window is displayed so the method and/or sample data can be stored. Then the Main window is displayed.

6.5 Data Manipulation

To display stored scan file(s):

- 1. Before displaying stored scans, clear the display by clicking on <a h
- 2. Verify that the desired "Overlay" selection is displayed.
- 3. To select the stored Scan file(s), click on "VIEW" following "Scan directory" to display the Scan Directory window.
- 4. Click on to highlight the desired file name(s), then click on [OK] to remove the directory window and display the scans. If "Overlay" is enabled, up to 12 file names can be selected. Once the scans are displayed, they can be manipulated using the instructions in this and the following section.
- 5. The scan speed is displayed following the scan file name on the Scan window.

The presentation of the spectral data can be optimized using the following features:

Ordinate label - The ordinate can be changed from absorbance to transmittance by clicking on the label.

Limit changes - The limits on the ordinate and/or wavelength axes can be changed by clicking on the displayed value and inputting the desired value.

Zoom - The "zoom" feature can be used to expand any portion of the graph. Click on **Zoom**, then click on two points on the graph to place crosses at the opposite corners of the area to be enlarged. When the second cross is clicked on, the graph is replotted. This can be repeated as often as desired. To return to the original plot, click on **ZoomOut**.

Trace - "Trace" can be used to display the ordinate value and the wavelength at any point on the plot. Click on <Trace>. Then move the arrow to the point of interest on the plot and click on the center mouse button to place a vertical line on the spectrum. The values at the place where the vertical line is placed are displayed in the upper right-hand side of the window. To move the vertical line to either the right or left, click on the right or left mouse button, respectively, with the arrow placed anywhere on the plot. To move to a different position on the spectrum, position the

Wavelength Scan 6-15

arrow and click on the center mouse button. An asterisk after the reading indicates that it is out of range.

NOTICE

The digital data at all wavelengths are displayed in the Tabulate window.

Autoscale - Automatically scale the ordinate axis. This command does not change the wavelength limits.

Annotate - The data can be annotated by clicking on <Annotate>. Then, click on the graph to position a cross and input information from the alphanumeric keypad or keyboard. Up to four annotations can be placed on the graph. The annotations are printed with the window, but are not stored with the data.

Smoothing - Selection of the number of data points to use for smoothing affects the peaks that are picked by the instrument. If too few points are used, extraneous or insignificant peaks are picked. If too many points are used, real peaks are flattened or lost. Click on the displayed smoothing value and input different values, visually examining the data to determine the optimum smoothing value.

6.6 Function Selection

The Function Selection window, Figure 6-5, is used to select derivative or log spectral display and to select peak, valley and/or point pick tabulation. It is displayed when **Function** is clicked on from the Wavelength Scan window, or "VIEW" from the Method window.

Wavel Print	ength '	Scan	•	Function Sele		kit,
	Scan First Secon Third Fourtl Log	Der d Der Der h Der	iva iva iva	ative vative ative vative		
	Peak I Number		P	aks: 5		
	Valle: Number			alleys: 5		
	Point	Pick	t			
Wavel	ength	Use	9	Wavelength	Use	
256 306 356 406).0).0).0).0).0	ENO ENO ENO ENO ENO ENO]	500.0 550.0 600.0 650.0 700.0 750.0	INO INO INO INO INO]

Figure 6-5. Function Selection Window

To select the desired function(s):

- 1. Verify that the data from the desired sample(s) are displayed on the Wavelength Scan window.
- 2. Click on <Function> to display the Function Selection window.
- 3. Click on to darken the box to select the desired function(s).

Derivative and log - Click on to select the desired information to be plotted for the selected scan file. If "Overlay" is disabled, any number of selections can be made. If "Overlay" is enabled, only one selection can be made.

Peak / Valley Pick - Click on to select the feature and input the number of peaks and/or valleys to pick.

Range: 1 to 30 Default: 5

Point Pick - Click on to select the feature. Click on the displayed wavelength value(s) to input the desired value(s) and click on to change the "Use" column to [Yes].

- 4. When the selections are made correctly, click on **Exit>** to remove the window from the display.
- 5. Example data, plotted with the selections in the Function Selection window (Figure 6-5), are shown in Figure 6-6.

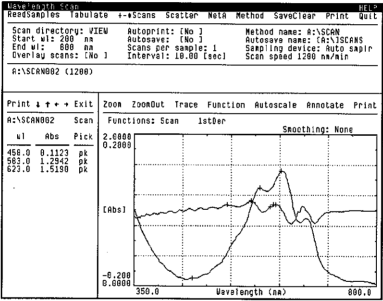


Figure 6-6. Wavelength Scan Window with Multiple Functions

The ordinate axis labels for the different selections are stacked on the lefthand side of the plot. The labels can be changed using the instructions in section 6.5.

The peak pick data for the absorbance scan is displayed. To display the peak pick data for the first or second derivative, click on <->>, located in the menu bar above the peak pick data.

6.7 Tabulated Data

To print the absorbance value and corresponding wavelength at each point in the spectrum, click on **Tabulate>**. The first 11 data points from a typical spectrum are shown in Figure 6-7.

Tabulat	ion for	Scan	File:	A:\SCAN001
wl	Abs			
200.0 201.0 202.0 203.0 204.0 205.0 206.0 207.0 208.0	0.1122 0.1036 0.0965 0.0904 0.0829 0.0757 0.0691 0.0636	3 5 4 9 7		
209.0 210.0	0.0526 0.0485	-		

Figure 6-7. Tabulated Scan Data

Wavelength Scan 6-19

6.8 Spectral Addition, Subtraction and Multiplication

Spectra that have been stored can be scaled using a factor, then can be added, subtracted or multiplied. The +-* Scans window, Figure 6-8, is used to select these calculations. It is displayed when <+-*Scans> is clicked on from the Wavelength Scan window.

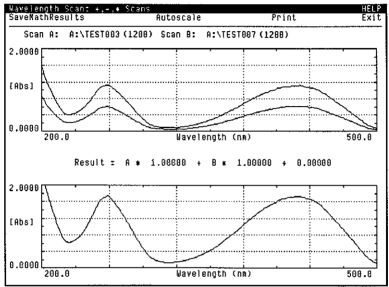


Figure 6-8. +-* Scans Window

The +-* Scans window has two graphic portions. The upper graph is used to display the raw data for the two spectra. The lower graph displays the resultant spectrum.

To perform the manipulation:

- 1. To input the file name for the first spectrum, A, click on "Scan A". The Scan Directory window is displayed. Click on the desired file, then [OK]. The spectrum is displayed.
- 2. Repeat step 1 to select the file for scan B.
- 3. The format of the spectra can be changed by clicking on the ordinate label and the ordinate and wavelength axis limits, and inputting the desired values.

4. To input the calculation parameters, click on "Result" in the middle of the window to display the +-* Parameters window, Figure 6-9.

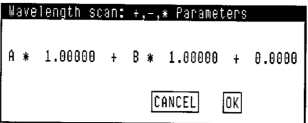


Figure 6-9. +-* Parameters Window

- 5. Three factors can be input in the window. These are a scaling factor for Scan A, a scaling factor for Scan B, and an additional value that is added to (subtracted from or multiplied by, depending upon sign) the results. Click on each of these factors and input the desired value.
- 6. Whether the spectra are added, subtracted or multiplied is determined by the sign. Click on to toggle between + (for addition), (for subtraction), and * (for multiplication).
- 7. When the desired parameters are displayed, click on [OK] to use the equation and remove the +-* Parameters window from the display.
- 8. The equation and the resultant spectrum are displayed in the lower graph. The format of the spectrum can be changed by clicking on the function, ordinate or wavelength axis limits and inputting the desired values.
- 9. To store the resultant spectrum, click on <SaveMathResults> to display the Save Results window. Input the desired file name, then click on [OK] to remove the window and store the data.
- 10. To print the window, click on <Print>.
- 11. To exit the +-* Scans window and return to the Wavelength Scan window, click on <Exit>.

6.9 Scatter Correction

The Scatter window, Figure 6-10, is used to correct a stored spectrum for scatter. It is displayed when **<Scatter>** is clicked on from the Wavelength Scan window.

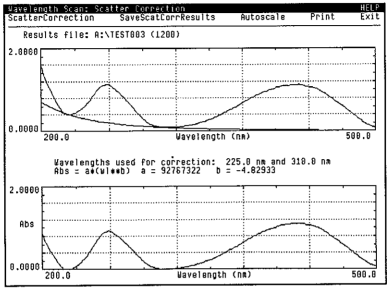


Figure 6-10. Scatter Window

The Scatter window has two graphic portions. The upper graph is used to display the uncorrected spectrum and the scatter component. The lower graph displays the spectrum corrected for scatter.

To correct for scatter:

- 1. Click on "Scan file". The Scan Directory window is displayed. Click on the desired file, then click on [OK]. The spectrum is displayed on the upper graph.
- 2. The format of the spectrum can be changed by clicking on the function, ordinate or wavelength axis limits and inputting the desired values.
- 3. Click on **<ScatterCorrection>**, then click on two points in the spectrum to place crosses where there is no sample absorption; that is, where all apparent absorption is due to scatter. When the second cross is clicked on, the calculation is performed. The scatter spectrum is displayed in the upper graph, the corrected sample spectrum is

displayed in the lower graph, and the equation for scatter correction is displayed in the middle of the window, along with the constants used to calculate the correction.

As an alternative, the wavelength values in the center of the window can be clicked on and new wavelength values input.

- 4. The format of the spectrum can be changed by clicking on the function, ordinate or wavelength axis limits and inputting the desired values.
- 5. To store the resultant spectrum, click on <SaveScatCorrResults> to display the Save Results window. Input the desired file name, then click on [OK] to remove the directory window and store the data.
- 6. To print the window, click on <Print>.
- 7. To exit the Scatter Correction window and return to the Wavelength Scan window, click on **Exit**.

6.10 Net Absorbance Calculations

The Net Absorbance Calculations window, Figure 6-11, is used to calculate net absorbance from a stored spectrum. Either one or two baseline points can be used. It is displayed when <NetA> is clicked on from the Wavelength Scan window.

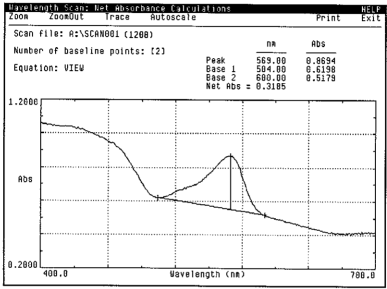


Figure 6-11. Net Absorbance Calculations Window

To calculate net absorbance:

- 1. Click on "Scan file". The Scan Directory window is displayed. Click on the desired file, then click on [OK]. The spectrum is displayed.
- The format of the spectrum can be changed by clicking on the absorbance or wavelength axis limits and inputting the desired values.
 Trace and <Zoom can also be used. Refer to section 6.5 for detailed instructions.
- 3. Select the desired number of baseline points by clicking on the value following "Number of baseline points" to change from [1] to [2].
- 4. If desired, the equation can be displayed by clicking on "VIEW" following "Equation".

- 5. The wavelength values for the peak and the baseline point(s) are displayed in a table at the top of the window. To input the desired wavelengths:
 - a. Click on "Peak", "Base 1", or "Base 2" to select the desired point.
 - b. Use the mouse buttons to position the "Trace" indicator at the desired wavelength on the graph:

Center mouse button - move to the data point closest to the position of the arrow.

Right mouse button - with the arrow positioned anywhere on the graph, move one data point to the right.

Left mouse button - with the arrow positioned anywhere on the graph, move one data point to the left.

The position of the "Trace" indicator line is shown in the table. As an alternative, the desired wavelength can be input in the table by clicking on the displayed wavelength value and inputting the desired wavelength value.

- c. Repeat steps a and b until the desired wavelengths are displayed.
- d. Click above the graph on a blank part of the display to remove the trace line and accept the displayed wavelengths. The net absorbance value is calculated, and the lines indicating the points used in the calculation are placed on the spectrum.
- 6. Examine the data to verify that the desired points were selected. If desired, change any of the points, using the instructions in step 5, above.
- 7. To print the window, click on **Print**.
- 8. To exit the Net Absorbance Calculations window and return to the Wavelength Scan window, click on **Exit**.

6.11 Example Analyses

EXAMPLE 1

Obtain a spectrum of a purified protein sample. Find the absorbance maxima and tabulate the absorbance data at 230, 260, 280, and 320 nm. Store the sample spectrum in file "A:\BSA001".

SOLUTION

Collect the data on the Wavelength Scan window, as shown in Figure 6-12. Use the Function window to select Peak Pick and input the four wavelengths for Point Pick. Be sure to change the "Use" column to [Yes]. Use << PrtScrn>> to print the window, so that the point pick data are on the same printout as the spectral data.

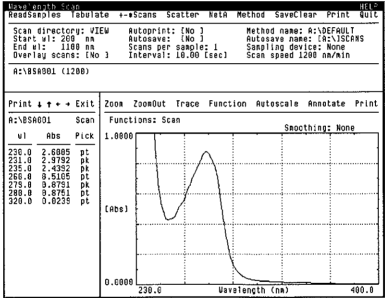


Figure 6-12. Sample Spectral Data with Point Pick

Display the spectrum shown in Example 1 with the first derivative overlaid.

SOLUTION

With the spectra displayed in Figure 6-12, click on **Function** to display the Function window. Click off "Peak Pick" and "Point Pick", and click on "I Der". The overlaid spectra are shown in Figure 6-13. The absorbance axis limits are 0.0 to 1.0. The first derivative axis limits are -0.5 to 0.5.

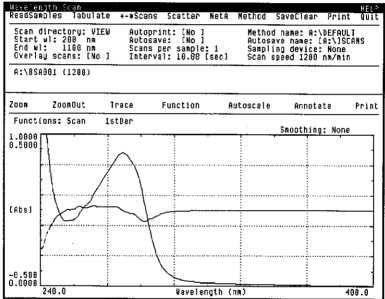


Figure 6-13. Overlaid Multifunction Spectra

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An analgesic sample contains 12.5 μ g/mL caffeine. Subtract the caffeine from the sample spectrum. The sample spectrum is file "A:\SAMP001". The caffeine spectrum (25 μ g/mL) is file "A:\CAFFN001".

SOLUTION

Use the +-* Window, shown in Figure 6-14. Input Scan A as "A:\SAMP001" and Scan B as "A:\CAFFN001". In the +-* Parameters window, input a factor of 1 for A, a factor of 0.5 for B, add a factor of 0, and set the first sign to "-".

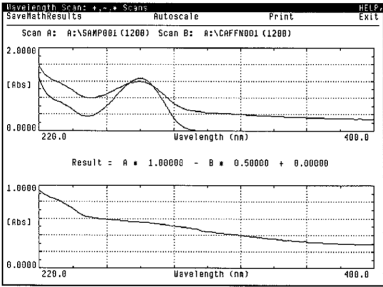


Figure 6-14. Component Subtraction

The resulting spectrum in Example 3 looks like scatter. Perform scatter correction on the sample spectrum.

SOLUTION

With the +-* Window displayed, click on **Exit** to display the Wavelength Scan window. Then click on **Scatter** to display the Scatter Correction window shown in Figure 6-15. Input the Scan file "A:\SAMP001". Click on **ScatterCorrection** and input wavelengths of 350 and 400 nm. The spectrum corrected for scatter is displayed on the lower part of the window.

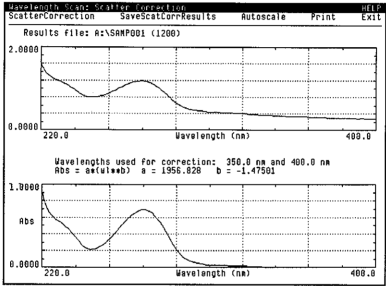


Figure 6-15. Caffeine Sample Corrected for Scatter

Wavelength Scan

6.12 Data Output

Spectral data are output to the Communications port whenever an Output Data Type is selected in the Communications Configuration window. The type of data output is selected: "raw and calculated", "raw", or "calculated". General information on data output is provided in section 9.3.

In the Wavelength Scan mode, the same data are sent, regardless of which output type is selected. The data output is the wavelength, followed by the reading in either absorbance or transmittance, for the entire wavelength range.

The following is the output obtained for a scan from 200 to 210 nm in absorbance. The "User ID" is input in the Communications Configuration window. The "Method ID" is the method name. A <a reaction and line feed> are sent at the end of every line.

User ID: DU600
Method ID: A:\SCAN
Date: 10\09\91
Time: 10:38
200.000 0.112
201.000 0.103
202.000 0.096
203.000 0.090
204.000 0.082
205.000 0.075
206.000 0.069
207.000 0.063
208.000 0.057
209.000 0.052
210.000 0.048

6.13 Files

Two file types are created in the Wavelength Scan mode: method files and scan files. The method files are stored in the SCAN_M directory with the .APX extension. Method files are ASCII files and cannot be converted to the Lotus format.

The scan files are stored in the SCAN_D directory with the .SCN extension. They contain scan parameters and data at every wavelength in the scanned range.

6.14 ASCII Format

The ASCII file for a wavelength scan consists of two parts: analysis parameters and sample data.

The sample data include the wavelength, followed by the reading in either absorbance or transmittance, for the entire wavelength range.

The following ASCII file was converted from an absorbance scan from 200 to 210 nm.

Start Wavele End Waveleng Read Mode: Read Average Nanometer In Time: 0	th: 210.0000 ABS : 0.1000 :terval: 1.0000
200.0000 201.0000 202.0000 203.0000 204.0000 205.0000 206.0000 207.0000 208.0000 209.0000 210.0000	0.1036 0.0965

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6.15 Lotus Format

The Lotus file for a wavelength scan consists of two parts: analysis parameters and sample data.

The sample data start in row 11 and are stored in the following columns:

Column A Wavelength

Column B Reading in either absorbance or transmittance

The following Lotus file was converted from an absorbance scan from 200 to 210 nm.

49.2	Start Waveleng	h: 200
- 2	End Wavelength:	210
3.	Read Mode:	ABS
- 4	Read Average:	0.1
- 5	Nanometer Inter	
. 6	Time:	g g
7		
9		
J. J. J.	A B	The same of the sa
-11	200 0.1	
12	201 0.10	36
13	202 0.09	55
	203 0.09	04
15	204 0.08	29
16	205 0.01	57
17	206 0.00	91
	207 0.00	31
-18		
18 19	208 0.09	74
18 19 20		

SECTION SEVEN

KINETICS/TIME

Kinetic data are collected in absorbance as a function of time. Multiple samples can be analyzed using an Auto Cell Holder, although the sampler is not required for kinetic analysis.

The data are plotted or tabulated as they are collected and are stored in the instrument memory. Single or multiple blanks can be specified. A second wavelength can be selected for background correction.

After data collection is complete, the user can calculate the rate of the kinetic reaction using a linear regression calculation over the desired time interval for the best fit straight line. Because the data are stored in memory, these calculations can be repeated until the optimum results are obtained.

7

7.1 Principles of Operation

Temperature Control

If the Temperature Controller is installed, temperature is controlled automatically in this mode. When <ReadSamples> is clicked on, the Temperature Controller is turned on and set to the input temperature. This allows time for the cell holder to equilibrate at the reaction temperature before [START] is clicked on. The cell holder can be preheated, if desired, using the Device Control window to turn on the Temperature Controller.

The cell holder changes temperature quickly. However, it takes about 10 minutes for the Temperature Controller to warm the solution in a standard cuvette from 25 to 37 degrees, because of slow heat transfer through the glass walls of the cuvette. Therefore, it is suggested that the solution be pre-warmed before being placed in the cell holder. The Temperature Controller can efficiently maintain the temperature of pre-warmed solutions.

Instrument Blank

The blank reading for the instrument should be taken on a cuvette filled with solvent (substrate) before the analysis begins. The solvent should be at the same temperature that the analysis will be performed.

Match Correction

When using an Auto Cell Holder, Match can be used to correct for differences in the cell positions by taking a zero reading at an input wavelength on each cell position (with a cuvette filled with blank solution and inserted into the Auto Cell Holder). When Match correction is enabled, all readings taken by the instrument will be corrected by the offsets displayed on the Match window. Match correction should be used with the Micro Auto-6 and the Micro Auto-12 Accessories.

Background Correction - Net Absorbance

It is possible to correct for raised or sloping baselines, which may be caused by turbidity in the sample, using background correction. The user can input one background wavelength for background correction. If a background wavelength is selected, the reading at the background wavelength is subtracted from the reading at the analytical wavelength to calculate a net absorbance reading, $A_{\rm na}$.

$$A_{na} = A_a - A_{bk}, \qquad (1)$$

where, A_a is the reading at the analytical wavelength and A_{bk} is the reading at the background wavelength. Match correction, if enabled, is performed on all cell positions prior to this calculation.

Blank Subtraction - Corrected Absorbance

Any number of blank(s) and sample(s), up to 12 total, can be analyzed and stored in the same Results file. The user specifies which are blanks and which are samples. If more than one blank is specified, the sample(s) that follow each blank will be corrected by the preceding blank.

When rate results are calculated, the blank solution reading is subtracted from the corresponding sample reading at each time point for every sample.

$$A = A_s - A_{bl}, (2)$$

where, A is the corrected absorbance, A_s is the sample reading and A_{bi} is the blank solution reading at each time point. Match correction and/or background correction calculations are performed on both blanks and samples prior to this calculation.

Concentration Calculation

The corrected absorbance (A, equation 2) can be multiplied by a factor to calculate concentration, using the equation:

$$Concentration = A x Factor$$
 (3)

Rate Calculation

The rate is calculated using linear regression to determine the best fit of a straight line to the data, using the basic rate equation:

Rate =
$$(A_2 - A_1) / (t_2 - t_1)$$
 (4)

where, the rate is calculated in units of delta A/minute, the corrected absorbance (equation 2) is used for each of the absorbance readings, and the limits of the time interval are selected by the user. The rate is calculated for samples, only. The calculated line is displayed with the actual data so that the user can compare the calculated rate to the actual data. The variance is also calculated and displayed with the rate. The variance gives the user an indication of how well the data fit the calculated line.

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Enzyme Activity Calculation

The rate can be multiplied by a user-input factor to convert it to enzyme activity or other concentration units.

Enzyme Activity = Rate x Factor.
$$(5)$$

Data Storage

A single Results file can hold data from up to 12 samples; the number is input by the response to the "number of samples" parameter. Each sample is designated as a sample or a blank, using the "sample assignment" information on the data collection window, or on the Method window.

It is necessary to store data from various samples in the same Results file, if the analysis of the data requires that the data from the samples be compared. Also, data from blanks must be stored in the same file as the associated sample data.

Samples that are analyzed singly are added to the Results file until the input "number of samples" is reached. Then the Results file is complete.

When the Auto Cell Holder is used, "groups" of sample data will be added to the Results file until the input "number of samples" is reached. The size of the "group" is equal to the "number of cells" parameter set for the Auto Cell Holder on the Sampling Device window.

For example, if the "number of cells" is 4 and the number of samples is 12, the Auto Cell Holder will be used three successive times to analyzed four samples. Then the Results file is complete.

7.2 Parameter Setup

Click on "KINETICS/TIME" from the Main window to enter the analysis mode. The Plotting window, Figure 7-1, is displayed. The Plotting window is used to setup analysis parameters and display kinetic data graphically as they are collected. (There is another data collection window in the Kinetics/Time mode - the Tabulation window.)

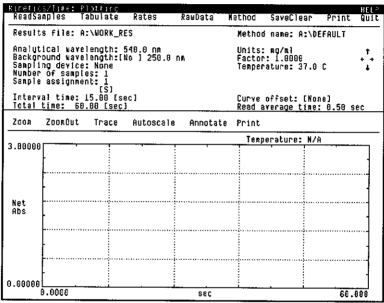


Figure 7-1. Plotting Window

Use the Method window to setup the analysis parameters:

- Click on <Method> to display the Method window, Figure 7-2. The
 Method window is used to setup analysis parameters, recall stored
 methods and create new methods. General information on method
 windows is provided in section 3.8.
- 2. To recall a stored method, click on the desired method name in the listing at the top of the Method window.

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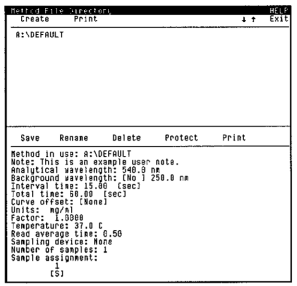


Figure 7-2. Method Window

3. The analysis parameters are displayed on the lower part of the Method window. Input the desired analysis parameters:

Method in use - This displays the name of the method that has been selected. If the method is protected, **PROTECTED** is displayed following the method name. If the method is protected, the analysis parameters cannot be changed. To input a new method name, click on <Create>.

Note - Click on to input a 40-character message that is used to described the method or procedure.

Analytical wavelength - The wavelength for data collection.

Background wavelength - The readings collected at the background wavelength are subtracted from the reading at the analytical wavelength to calculate the net absorbance. Toggle between [Yes] and [No] to enable or disable the correction.

Interval time - The time between successive readings of the same sample. The minimum time is affected by the number of cells and the read average time.

Using a read average time of 0.5 second, the following are the minimum interval times in seconds, based upon the type of Auto Cell Holder and the number of cells.

Auto 6 Cell Holder		Aut	Auto 12 Cell Holder						
Cells	Time	Cells	Time	Cells	Time				
1	0.05*	1	0.05*	7	9.9				
2	3.1	2	2.9	8	11.4				
3	4.8	3	4.3	9	12.8				
4	6.6	4	5.7	10	14.1				
5	8.3	5	7.2	11	15.7				
6	10.1	6	8.3	12	17.2				

A read average time of 0.05 second was used.

Total time - The instrument collects data until the total time has elapsed.

Curve offset - This parameter will automatically offset the first reading on each cell, so that the plotted data will not overlay. Toggle between [Ascending], [Descending] and [No] to enable or disable the offset. [Ascending] is for increasing reactions and [Descending] is for decreasing reactions.

Units - Input up to an 8-digit alphanumeric name for the enzyme activity units.

Factor - Input a factor to convert the absorbance to concentration units and the rate to enzyme activity.

Temperature - Input the operating temperature for the Temperature Controller, if used.

Read average time - The time, in seconds, that data are collected and averaged to take a reading. Ten sets of data are collected every second. If the interval time is less than 1 second, a read average time of 0.1 is always used.

Sampling device - Display the Sampling Device window for selection of the sampling device.

Number of samples - The number of sample data to be stored in a single Results file. If using the Auto Cell Holder, the "number of cells" input on the Sampling Device window cannot be greater than the input to this parameter. If it is greater, then the input to this parameter is used for the "number of cells".

Sample assignment - Click on to identify the samples stored in a single Results file. [B] is for blank; [S] is for sample. The sample(s) following each blank will be corrected for that blank reading.

- To store the analysis parameters in the selected method file, click on Save>.
- 4. Click on **<Exit>** to display the Plotting window with the parameters from the selected method.

To input the desired parameters on the Plotting window:

- 1. The analysis parameters are listed near the top of the window. To change any of these, click on the parameter and input the desired value. A description of the parameters is provided above.
- 2. The absorbance limits are input by clicking on the displayed values on the graph and inputting the desired values.

7.3 Data Collection

The data can either be plotted or tabulated as they are collected. If the Plotting window is displayed and tabulation is desired, click on <Tabulate>. If the Tabulation window is displayed and a plot is desired, click on <Plot>. Either the Plotting or the Tabulation window can be used for data collection.

To collect data using the displayed analysis parameters:

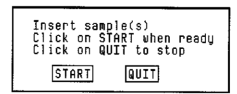
1. Place a cuvette of substrate in the cell holder. Click on <<BLANK>>. The instrument blanks on the substrate.

<u>Auto Cell Holder</u> - Place a cuvette of substrate in the cell position that is in the light beam, then click on <<BLANK>>. The instrument blanks on the substrate.

2. If using the Auto Cell Holder, Match can be used to correct for slight differences in the cuvettes. Click on <<MATCH>> to display the Match window. Place a cuvette filled with substrate in each of the cell positions that will be occupied with a blank or sample. Take the readings and enable Match correction.

The cuvettes should not be removed from the instrument before the sample analysis. If the cuvettes are removed, it is necessary to return each cuvette to the same cell position in the same orientation. For most accurate results, Match correction should be repeated each time the cuvettes are removed from the instrument.

- 3. If the Plotting window is displayed, set the limits on the absorbance axis by clicking on the displayed value and inputting the desired value.
- Click on <ReadSamples>. The Read Samples window is displayed, which is used to read the samples.



The Temperature Controller turns on, if not already on, and the temperature is controlled to the input temperature parameter. Wait

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until the temperature display on the Temperature Controller reaches the desired temperature. (The actual temperature is placed on the Plotting window when [START] is pressed.)

5. Place a cuvette of sample in the cell holder and click on [START]. The data are plotted or tabulated as they are collected, depending upon the window.

<u>Auto Cell Holder</u> - Load the blank(s) and samples in the order indicated on the top of the window. Positions are indicated with a "B" for blank and a "S" for sample. Click on [START].

- 6. The analysis can be terminated at any time by clicking on [QUIT]. However, after [QUIT] is clicked on, no additional data can be added to the Results file, even if all the samples have not been analyzed.
- 7. When data collection is complete, if the total "number of samples" has not been analyzed, the window shown in step 4 is displayed. Repeat step 5, until all samples have been analyzed.

NOTICE

Do not change from the Plotting to Tabulation window, or display the Rates window until all samples have been analyzed.

8. When all the samples input for "number of samples" have been analyzed, the window shown in step 4 is removed. The data collection window can be changed from Plotting to Tabulation by clicking on <Tabulate>, or from Tabulation to Plotting by clicking on <Plot>. A typical Plotting window is shown in Figure 7-3. A typical Tabulation window is shown in Figure 7-4.

The data on the Plotting window can be reformatted and annotated. Detailed instructions are provided in the next section.

The net absorbance readings (equation 1) are tabulated on the Tabulation window. These readings have been corrected for background absorbance and Match correction, if used. All the raw data, plus additional calculated data are displayed on the Raw Data window.

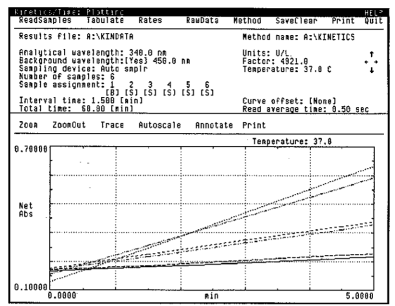


Figure 7-3. Plotting Window

	-		_			
Kiretics/Tir ReadSamples	e: Tabulation Plot	Rates	RawData	Method	SaveClear	HEL⊃ Print Quit
Results fil	e: A:\KINDATA	~~~		Method	name: A:\KI	INETICS
Background Sampling de Number of s		s] 450.9 n lr			ህ/L : 4921.0 ature: 37.0	c
Total time:	me: 1.580 (min 60.00 (min)	(\$) (\$) (\$1 [\$]		offset: [No verage time:	
Net #bsorbar Cell #		2	1		4	
Sample ID Time min	1 1	2	3		4 ! 4 !	5 6 5 6
0.880 1.500 3.080 4.500 6.000 7.500 10.500 12.000 15.000 16.000 16.000 18.000 19.500 21.000	0.1905 0.1965 0.2149 0.2248 0.2498 0.2679 0.3841 0.3201 0.3455 0.3566 0.3774 0.3935 0.4110 0.4483	0.1057 0.2395 0.2395 0.3544 0.4144 0.4729 0.5306 0.6402 0.7064 0.7656 0.87723 0.87723 0.9314 1.0519	0.1650 0.2867 0.4065 6.5286 6.6517 6.8980 1.0502 1.2683 1.38943 1.38943 1.7529 1.77520 1.8721 2.6697	0.19; 0.24; 0.31; 0.36; 0.49; 0.55; 0.61; 0.67; 0.79; 0.85; 0.95; 1.04;	96 6.27 98 9.42 92 6.57 91 9.86 937 1.86 96 1.15 96 1.29 98 1.59 97 1.89 97 2.082 18 2.17	74 0.2023 27 0.2245 80 0.2470 84 0.2675 49 0.2911 82 0.3094 80 0.3296 80 0.3749 80 0.3749 81 0.4377 81 0.4577 81 0.4589

Figure 7-4. Tabulation Window

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- 9. The rates of the kinetic reactions are calculated using the Rates window, described in section 7.5.
- 10. When all data collection is complete, use the Device Control window to turn off the Temperature Controller. The fan will turn off about 5 minutes after the Temperature Controller is turned off.
- 11. To clear all sample data from the window or to store the data, then clear the window, click on <SaveClear>. The Save Clear window is displayed so the sample data can be stored. If the data are stored, no additional data can be placed in the Results file, even if all the samples have not been analyzed.

NOTICE

Results files can be recalled in the Enzyme Mechanism mode, if they are copied into the "MECH_D" directory. The File Utilities mode is used to copy files.

12. When the analysis is complete, click on <Quit>. The Quit window is displayed so the method and/or sample data can be stored. Then the Main window is displayed.

7.4 Plotted Data

The presentation of plotted data on the Plotting and the Rates windows can be optimized using the following features:

Limit changes - The limits on the absorbance and/or time axis can be changed by clicking on the displayed value and inputting the desired value.

Curves offset - If the plotted data for several samples overlay at the zero time point, the data can be offset by enabling the "Curves offset" parameter.

Zoom - The "zoom" feature can be used to expand any portion of the graph. Click on <Zoom>, then click on two points on the graph to place crosses at the opposite corners of the area to be enlarged. When the second cross is clicked on, the graph is replotted. This can be repeated as often as desired. To return to the original plot, click on <ZoomOut>.

Trace - "Trace" can be used to display the time at any desired position on the plot. Click on <Trace>. Then move the arrow to the point of interest in the plot and click on the center mouse button to place a vertical line on the plot. The values at the place where the vertical line is placed are displayed. To move the vertical line to either the right or left, click on the right or left mouse button, respectively, with the arrow placed anywhere on the plot. To move to a different position on the plot, position the arrow and click on the center mouse button. An asterisk after the reading indicates that it is out of range.

NOTICE

The digital data at all wavelengths are displayed on the Raw Data window.

Autoscale - Automatically scale the ordinate axis. This command does not change the time limits.

Annotate - The data can be annotated by clicking on <Annotate>. Then, click on the graph to position a cross and input information from the alphanumeric keypad or keyboard. Up to four annotations can be placed on the graph. The annotations are printed with the window, but are not stored with the data.

7.5 Rate Calculation

The Rates window is used for rate calculation. It displays the plotted data, corrected for the blank, and allows the user to input the necessary information to calculate the reaction rate between any two time points on the graph.

The Rates window, Figure 7-5, is displayed when <Rates> is clicked on from either of the data collection windows. The Results file cannot be changed on the Rates window, so verify that the desired Results file is displayed on the data collection window before displaying the Rates window. Information about the data in the selected Results file is displayed in the upper portion of the Rates window.

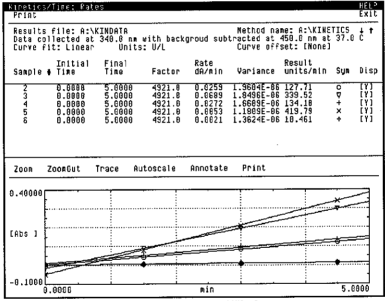


Figure 7-5. Rates Window

To calculate the rate of the kinetic reaction using the Rates window:

1. All sample data stored in the Results file are plotted. (Blanks are subtracted from the corresponding samples and not plotted.) Plotted data for any sample(s) can be deleted from the display by clicking on the [Y] in the "Disp" column in the table, replacing it with an [N] for "no".

- 2. The presentation of the plotted data can be optimized by using the curves offset, by using "zoom" or by changing the limits individually. Detailed instructions are provided in section 7.4.
- 3. The label on the ordinate axis can be changed from [Abs] to [Conc]. Concentration is calculated by multiplying the factor by each absorbance reading (equation 3).
- 4. The rates are calculated using the parameters input in the "Initial Time", "Final Time", and "Factor" columns. The time points can be found from the plotted data using Trace.

To input the same value for all cells, click on the column heading. Input the desired value in the displayed window and click on [OK]. To input different values for each cell, click on displayed value and input the desired value.

The line displayed on the graph is the slope of the line calculated from the initial and final time points. These times can be changed to optimize the fit of the calculated line to the collected data.

- 5. The tabulated results include the rate in delta A per minute (equation 4), the variance calculated for the curve fit, and the enzyme activity in the input units calculated from the factor (equation 5).
- 6. The table of rate results is printed by clicking on <**Print>**, located in the menu bar above the table. The plotted data are printed by clicking on <**Print>**, located in the menu bar above the graph. The entire display is printed by clicking on <**PrtScrn>>**.
- 7. When the rates are calculated, click on **Exit** to return to the Plotting window.

7.6 Raw Data

The Raw Data window, Figure 7-6, tabulates the raw data at the analytical and background wavelengths, the net absorbance (background correction), and the corrected absorbance (blank subtraction). This window is displayed when **<RawData>** is clicked on from either of the data collection windows.

The Results file name cannot be changed on the Raw Data window, so verify that the desired Results file is displayed on the data collection window before displaying the Raw Data window. Information about the data in the selected Results file is displayed in the upper portion of the Raw Data window.

Biretin Print	s/Tiπe:	Raw data					(it
Result	s file:	A:\KINDATA			Method name: A:\KINETICS Data collected at 37.0	t	t
Sample	e assign	ment: 1 2 [8] [8]	3 4 [\$] [\$] [5 8 \$) [\$]			
Time (min)	Sample ID	Analyt wl 340.00nm	8kg w1 450.00nm	Net abs	Corrected Abs		
1.500	123456123456123456123456	6.2952 6.2551 6.4249 6.3889 6.4837 6.3878 6.3878 6.3147 6.5379 6.4364	0.1147 0.0734 0.2599 0.1882 0.3498 0.1218 0.1218 0.0751 0.2511 0.2511 0.3506	8.1865 9.1857 9.1657 9.1927 9.1349 9.1852 9.1963 9.2395 9.2496 9.2774	0.0052 -0.0155 0.0122 -0.0465 0.0047 0.0432 0.0904 0.0532 0.0818		
3.008	6 1 2 3 4 5	0.3273 0.3268 0.3701 0.6602 0.4978 0.7724	0.1250 0.1119 0.0786 0.2537 0.1870 0.3497	0.2023 0.2149 0.2995 0.4065 0.3108 0.4227	0.0060 0.0046 0.1916 0.0959 0.2078		
4.500	1 2 3 4 5 6	0.3471 0.3451 0.4289 0.7794 0.5571 0.9107	8.1226 9.1123 9.0749 9.2508 9.1908 9.3481 9.1202	0.2245 0.2328 0.3540 0.5286 0.3672 0.5706 0.2470	6.8096 6.1212 6.2558 6.1343 6.3378 6.0141		

Figure 7-6. Raw Data Window

To display the raw data and associated information:

- 1. The cell assignments for the Results file is displayed; they can be changed.
- 2. Only the first few readings are displayed. The other readings can be displayed by scrolling through the data, using the up and down arrows located in the upper right-hand corner of the window.

- 3. The tabular data include the net absorbance (equation 1) and the corrected absorbance (equation 2).
- 4. To print the window, click on <Print>. All data, even that which is not displayed because of insufficient room on the display, are printed on the printer.
- 5. When the Raw Data display is complete, click on **<Exit>** to return to the Plotting window.

7.7 Example Analyses

EXAMPLE 1

Setup the parameters for a kinetic analysis at 340 nm at 37°C. Analyze six samples, each with its own blank, using the Auto 6 Cell Holder. Take readings every 30 seconds for 5 minutes. Store the data in a file named "A:\CK_DATA".

SOLUTION

The analysis parameters are listed on the top of the Tabulation window, shown in Figure 7-7. The Auto 6 Cell Holder must be used twice to analyze all the samples. The data from the first three samples (with blanks) are tabulated on the window. The remainder of the samples will be analyzed subsequently, and stored in the same Results file.

Ripetics/Tim ReadSamples	e: Tapulation Plot	Rates Ra	awData	Method Sa	veClear (HELP Print Quit
	e: A:\CK_DATA				me: A:\CKT	EST
Background	wavelength: 34 wavelength:[Na vice: Auto sng annles: 12	1 450.0 nm		Units: U/ Factor: 4 Temperatu		• †
Sample assi Interval ti Total time:	grment: 1 2 [B] [S] me: 30.00 [sec 5.000 [min]	3 4 5 [B] [S] [B]	6 7 [S] [B]	Curve off	0 11 12] [B] [S] set: [None: age time: [
Net Absorbar	de Readings					
Cell # Sample ID Time sec	1 1	2 2	3	4 4	5 5	6
8,900 30,000 69,800 98,800 128,800 156,900 210,000 240,000 270,000 300,800	0.2139 0.2168 0.2168 0.2161 0.2107 0.2103 0.2113 0.2106 0.2106 0.2109 0.2113	0.2457 6.2451 6.2455 6.2458 6.2493 6.2525 6.2544 8.2578 6.2664 6.2663	0.2283 0.2288 0.2281 0.2281 0.2297 0.2292 0.2294 0.2294 0.2294 0.2299	0.2459 0.2429 0.2436 0.2436 0.2470 0.2532 0.2552 0.2558 0.2618 0.2644 0.2662	9.2197 6.2194 6.2194 6.2190 6.2196 6.2198 6.2192 9.2192 9.2192 9.2195 8.2194	0.2558 6.2567 9.2667 8.2654 6.2750 6.2805 9.2805 9.2911 9.2979 9.3035

Figure 7-7. Example Tabulation Window

After all the samples are analyzed, calculate the rate for the samples collected in Example 1, using the readings between 1 and 5 minutes. Use a factor of 4921 to calculate enzyme activity in units of U/L.

SOLUTION

With the Tabulation window displayed, click on <Rates> to display the Rates window, shown in Figure 7-8. Click on the headings for initial time, final time and factor and input the values of 1, 5, and 4921, respectively, to calculate the rates.

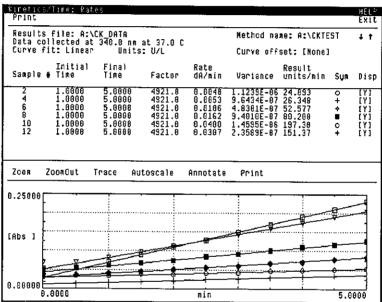


Figure 7-8. Example Rates Window

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Setup the parameters for a kinetic analysis of one sample at 37°. Read at 340 nm and correct for background at 500 nm. Take readings as frequently as possible for 30 seconds. Store the data in a file named "B:\CK FAST".

SOLUTION

The parameters are listed in the top of the Plotting window, shown in Figure 7-9. The data collected from the sample are plotted in the lower part of the window.

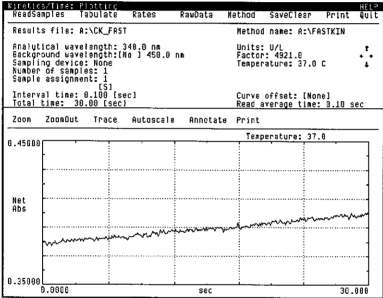


Figure 7-9. Example Plotting Window

7.8 Data Output

Data are output to the Communications port whenever an Output Data Type is selected in the Communications Configuration window. The type of data output is selected: "raw and calculated", "raw", or "calculated". General information on data output is provided in section 9.3.

In the Kinetics/Time mode, the order of the output for each data type is shown below. The "User ID" is input in the Communications Configuration window. The "Method ID" is the method name. A < carriage return > and < line feed > are sent at the end of every line.

Sample Data

If "raw and calculated" or "raw" is selected, the data output for each sample includes: sample number, time, and net absorbance. The following is the output obtained at the first two time points when the data in Figures 7-3 and 7-4 were collected. If "calculated" is selected, no information is sent.

User ID: DU600 Method ID: A:\KINETICS Date: 10/09/91 Time: 11:08 0.000 0.1805 0.000 0.1857 3 0.000 0.1650 0.000 0.1927 5 0.000 0.1340 0.000 0.1852 1.500 0.1963 1.500 0.2395 1.500 0.2867 1.500 0.2496 1.500 0.2774 1.500 0.2023

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Calculated Data at <Quit> or <SaveClear>

If either "raw and calculated" or "calculated" is selected, the rates information is sent each time the Quit window or the Save Clear window is removed from the display. If "raw" is selected, no information is sent.

The data output for each sample includes: sample number, initial time, final time, factor, rate, variance, and result. No data are sent for blanks. The following is the output obtained for the samples analyzed in the Rates window in Figure 7-5.

ı						
lle	er ID: D	HANN				
		A:\KINET	tce			
			100			
Da:	te: 10/0	9/91				
Ti	me: 11:0	8				
2	0.0000	5.0000	4921.0000	0.0259	1.9604E-06	127.712
3	0.0000	5.0000	4921.0000	0.0689	1.8496E-06	339.527
4	0.0000	5.0000	4921.0000	0.0272	1.6689E-06	134.180
5	0.0000	5.0000	4921.0000	0.0853	1.1809E-06	419.796
6	0.0000	5.0000	4921.0000	0.0021	1.3624E-06	10.4610

7.9 Files

Two file types are created in the Kinetics/Time mode: method files and data files. The method files are stored in the KIN_METH directory with the .APX extension. Method files are ASCII files and cannot be converted to the Lotus format.

The data files are stored in the KIN_DATA directory with the .DUF extension. They contain analysis parameters, rate data and sample data.

7.10 ASCII Format

The ASCII file for kinetics data consists of three parts: analysis parameters, rate data and sample data.

The rate data include: sample number, initial time, final time, factor, rate, variance and result. This is in the same format as the Rates window.

The sample data include the time, followed by: sample number, analytical absorbance, background absorbance (if enabled), net absorbance and corrected absorbance (samples, only.) This is in the same format as the Raw Data window.

The following ASCII file was converted from the data displayed in Figures 7-3 and 7-4. It includes the readings at the first two time points, only. The corresponding Rates window is Figure 7-5; the corresponding Raw Data window is Figure 7-6.

Background Background Number of S Sample assi	gnment:			s		
Interval ti		.5000 mi	n			
Total Time:		00 min				
	U/L					
	4921.0000					
Temperature Curve offse		JU0				
Read averag		0.5000				
kead averag	e time;	0.5000				
1	0.0000	60.0000	4921.0000	0.0111	0.0001	F. (07.
	0.0000	60.0000	4921.0000	0.0369	0.0001	54.4234
2 3	0.0000	60.0000	4921.0000	0.0652	0.1016	181.5955 320.6385
4 5	0.0000	60.0000	4921,0000	0.0388	0.0007	191,1480
5	0.0000	60.0000	4921.0000	0.0671	0.2096	330,3846
6	0.0000	60.0000	4921.0000	0.0138	0.0000	67.7595
0.0000	1	0.2952	0.1147	0.1805	0.1805	
	2	0.2591	0.0734	0.1857	0.1857	
	5	0.4249	0.2599	0.1650	0.1650	
	4	0.3809	0.1882	0.1927	0.1927	
	2	0.4837	0.3498	0.1340	0.1340	
1,5000	1	0.3070 0.3093	0.1218	0.1852	0.1852	
1.5000	,	0.3093	0.1129 0.0751	0.1963	0.1963	
	2 3 4 5 6 1 2 3	0.5379	0.0751	0.2395 0.2867	0.2395	
	4	0.4364	0.1869	0.2496	0.2867 0.2496	
	5	0.6280	0.3506	0.2774	0.2496	
	6	0.3273	0.1250	0.2023	0.2023	
			211230	012023	V.2023	

Kinetics/Time

7.11 Lotus Format

The Lotus file for kinetics data consists of three parts: analysis parameters, rate data and sample data. The rate data are in the same format as the Rates window. The sample data are in the same format as the Raw Data window.

The rate data start in row 18 and are stored in the following columns:

Column A	Sample number
Column B	Initial time
Column C	Final time
Column D	Factor
Column E	Rate
Column F	Variance
Column G	Result

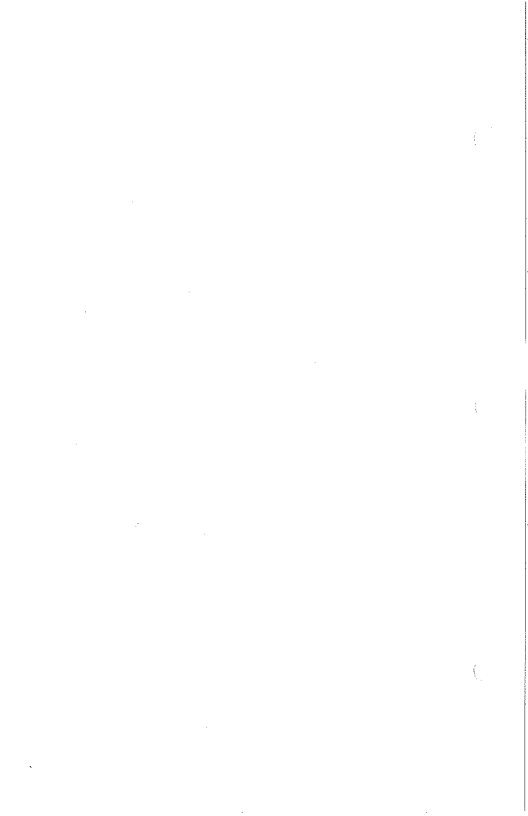
The sample data follow the rate data and are stored in the following columns:

Column A	Time data were conected
Column B	Sample number
Column C	Analytical absorbance
Column D	Background absorbance
Column E	Net absorbance
Column F	Corrected absorbance (samples, only.)

The Lotus file on the next page was converted from the data displayed in Figures 7-3 and 7-4. It includes the readings at the first two time points, only. The corresponding Rates window is Figure 7-5; the corresponding Raw Data window is Figure 7-6.

A		В	l e l	D L	E	P P	g
Analytical w		340					
Background w	ave	450					
Background c	orrYes						
Number of Sa		6					
Sample assig	nmeS	S	s	5	s	s	
Interval tim	e:	1.5 mi	n.			_	
Total Time:		60 mi	n				
Units:	U/L						
Factor:		4921					
Temperature:		37					
Curve offset		<u> </u>					
Read average	ti	0.5					
	9		D.	E		di di	
1	0	60		0.011059			
2	0	60		0.036902			
3	o	60		0.065157	0.101634	320.6384	
4	0	60		0.038843			
5	0	60		0.067137			
6	0	60	4921	0.013769	0.000014	67.75947	
0	1 0	.295150	0.114671	0.180478	0.180478		
	20	.259092	0.073417	0.185674	0.185674		
	3 0	.424870	0.259889	0.164981	0.164981		
	4 0	.380856	0.188184	0.192671	0.192671		
	5 0	.483736	0.349766	0.133970	0.133970		
			0.121845				
1.5			0.112927				
			0.075137				
			0.251149				
			0.186863				
			0.350603				
			0.124955				
			*********	0.202313	0.202313		

Kinetics/Time



The File Utilities mode is used to rename, copy, move and delete files, to provide disk status, to format new floppy diskettes, to convert files stored in the instrument to Lotus¹ or ASCII format, to receive files from and send files to an external computer.

The DU Series 600 Spectrophotometer is supplied with internal RAM memory, which is used for storage. This location is called drive A. An optional 1 MEG RAM can be added to this memory in drive A. Drive B is an optional 3½ inch diskette drive, which will accept 720 kB diskettes.

NOTICE

As an alternative, DOS commands can be used on an IBM compatible computer to rename, copy, and delete files stored on diskettes generated by the DU Series 600 Spectrophotometer.

¹Lotus is a trademark of Lotus Development Corporation.

8.1 File Utilities Window

The File Utilities window, Figure 8-1, is displayed when "FILE UTILITIES" is selected from the Main window.

List of Directories: A:\FIX METH B:\FIX METH A:\FIX DATA B:\FIX DATA A:\FIX DATA B:\FIX DATA A:\FIX DATA B:\FIX DATA A:\SCAN M B:\SCAN M A:\SCAN M B:\SCAN M A:\SCAN M B:\SCAN M A:\SCAN M B:\SCAN DATA A:\FIN DATA B:\FIN DAT

Figure 8-1. File Utilities Window

The menu bar commands for the File Utilities window are used to perform the various actions on a file. The instructions for each of these commands are given in the following sections.

The left-hand side of the window is used to display all the directories in the instrument. If optional application modes are installed, it may be necessary to scroll to display all the directories. The arrows in the upper right-hand corner of the window are used to scroll. When one of the directories is clicked on, the files in that directory are listed in the right-hand side of the window.

The desired file is selected from the file listing by clicking on the file name to highlight it. Clicking on a highlighted file name will remove the highlight. If the file listing fills the window, additional files can be displayed using the up/down arrows at the top right-hand corner of the window to scroll through the files.

8.2 Directories

The directories are created by the instrument as part of the format procedure (section 8.8) and are transparent to the user, except in the File Utilities mode. The user can neither create nor delete directories from the instrument.

Refer to the directory listing in Figure 8-1. Notice that each directory has a two-part name. The first part of the name identifies the operating mode; the second part of the name identifies the type of data. Also notice that there are identical directories for both drives A and B.

NOTICE

The instrument cannot read information stored in directories with names other than the ones listed.

The following tables explain the mnemonics used in the nomenclature.

File Utilities 8-3

Mnemonic	Mode
FIX	Fixed Wavelength
SCAN	Wavelength Scanning
KIN	Kinetics/Time
PROT	Protein Analysis
FRC	Fraction Read
NUC	Nucleic Acid
SCA	Single Component Analysis
ACT	Enzyme Activity
MECH	Enzyme Mechanism
MCA	Multicomponent Analysis
PROGRAM	Program mode - programs
PROGDATA	Program mode - data files
CUST_APP	Programs/Methods displayed on the Main window
LOTUS	Files converted to Lotus
ASCII	Files converted to ASCII
GELAR	Gel Scan Area
GELMW	Gel Scan Molecular Weights
GEL_SCNS	Gel data from either mode

Table 8-1. Directory Nomenclature - Mode Identifier

Mnemonic	Data Type	
METH or M	Method	
DATA or D	Data	
STDS or S	Standards	
SCNS	Scans	
CTRL	Controls	

Table 8-2. Directory Nomenclature - Data Type

8.3 Rename Command

To rename a file:

- Display the File Utilities window by clicking on "File Utilities" from the Main window.
- Click on the desired directory, listed on the left-hand side of the window. All the files in the selected directory are listed on the righthand side of the window.
- 3. Click on the desired file to highlight it.
- Click on <Rename>. The Rename File window, Figure 8-2, is displayed.

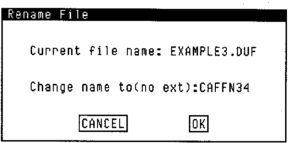


Figure 8-2. Rename File Window

- 5. The file name that was highlighted on the File Utilities window is displayed following "Current file name:".
- 6. Click on the file name following "Change name to(no ext):" and input the new file name using the alphanumeric keyboard. The instrument will automatically append the file name with the extension, so the file name must be input without the extension.
- 7. Click on [OK] to rename the selected file and remove the Rename File window from the display.

File Utilities 8-5

8.4 Copy Command

To copy a file from one directory to the same or other directory without changing the file name:

NOTICE

A file can be copied into any directory. However, the instrument is not able to read a file that is recalled in a different directory from which it was created. There are a few exceptions, which are noted in the respective operating mode section of this manual.

- 1. Display the File Utilities window by clicking on "File Utilities" from the Main window.
- Click on the desired directory, listed on the left-hand side of the window. All the files in the selected directory are listed on the righthand side of the window.
- 3. Click on one or more file names to highlight them.
- 4. Click on **<Copy>**. One of two Copy File windows is displayed, depending upon whether one or multiple files were highlighted.
- 5a. If one file was highlighted, the Copy File window, Figure 8-3, is displayed. Input the drive, directory and file name where the file is to be copied. Then click on [OK] to copy the selected file and remove the Copy File window from the display.

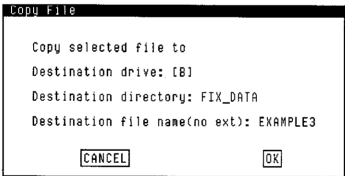


Figure 8-3. Copy File Window

5b. If multiple files were highlighted, the Copy Files window, Figure 8-4, is displayed. Input the drive and directory where the files are to be copied. The same file names are used. Then click on [OK] to copy the selected files and remove the Copy Files window from the display.

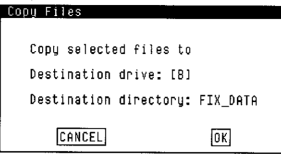


Figure 8-4. Copy Files Window

8.5 Move Command

To move one or more files from one directory to another without changing the file name(s):

NOTICE

A file can be moved into any directory. However, the instrument is not able to read a file that is recalled in a different directory from which it was created. A few exceptions are noted in the respective operating mode section of this manual.

- Display the File Utilities window by clicking on "File Utilities" from the Main window.
- Click on the desired directory, listed on the left-hand side of the window. All the files in the selected directory are listed on the righthand side of the window.
- 3. Click on one or more file names to highlight them.
- 4. Click on <Move>. The Move File(s) window, Figure 8-5, is displayed. Input the drive and directory where the file(s) are to be moved. Then click on [OK] to move the selected file(s) and remove the Move Files(s) window from the display.

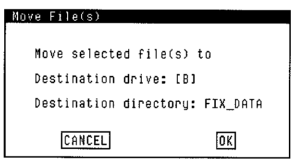


Figure 8-5. Move File(s) Window

8.6 Delete Command

To delete one or more files:

- 1. Display the File Utilities window by clicking on "File Utilities" from the Main window.
- 2. Click on the desired directory, listed on the left-hand side of the window. All the files in the selected directory are listed on the right-hand side of the window.
- 3. Click on one or more file names to highlight them.
- 4. Click on **Delete**. The Delete window, Figure 8-6, is displayed. Click on **[OK]** to delete the selected file(s) and remove the Delete window from the display.

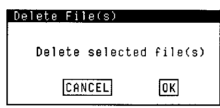


Figure 8-6. Delete File(s) Window

File Utilities 8-9

8.7 Disk Status Command

To display the amount of available space on each device:

- 1. Display the File Utilities window by clicking on "File Utilities" from the Main window.
- 2. Click on **<DiskStatus>**. The Disk Status window, Figure 8-7, is displayed. After reviewing the information, click on **[OK]** to remove the window from the display.

sk Statu	S	
Device	Capacity(bytes)	Free Space(bytes)
A B	1031936 730112	1020416 574464
		OK

Figure 8-7. Disk Status Window

8.8 Format Command

The Format command formats the internal memory (Virtual Disk A) or a 3½ diskette using the standard DOS procedure and sets up the directories for subsequent file storage. All diskettes used with the DU-600 should be formatted using this command.

NOTICE

A diskette can be formatted on an external computer, using the DOS format command. After the diskette is formatted, the appropriate DU-600 directories (defined in section 8.2) must be added to the diskette before it can be used with the instrument.

The format a diskette:

- Display the File Utilities window by clicking on "File Utilities" from the Main window.
- 2. Click on **<Format>**. The Format Disk window, Figure 8-8, is displayed.

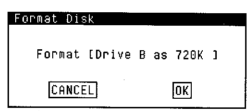


Figure 8-8. Format Disk Window

3. Verify that the proper drive and diskette size is displayed. The options are: [Virtual Disk A] and [Drive B as 720K]. Click on the displayed value to change it.

4. Click on [OK]. The following warning window is displayed.

WARNING
ALL DISK B CONTENTS WILL BE ERASED

Insert a new Diskette in Drive B

CANCEL OK

5. Insert the diskette into the drive B. Click on [OK] to format the diskette. While the diskette is formatting, a window with the cylinder and head being formatted is displayed. When the formatting is complete, the directories are set up, then the window is removed. It takes about 3 minutes to format a diskette.

8.9 Convert Command

Files converted to Lotus format are stored in the LOTUS directory. Files converted to ASCII are stored in the ASCII directory. Files in these directories cannot be recalled by the instrument. The format of the data stored in these files is provided at the end of the appropriate operating mode section in this manual or an accessory manual.

To convert one or more files to Lotus or ASCII without changing file names:

- 1. Display the File Utilities window by clicking on "File Utilities" from the Main window.
- Click on the desired directory, listed on the left-hand side of the window. All the files in the selected directory are listed on the righthand side of the window.
- 3. Click on one or more file names to highlight them.
- 4. Click on **<Convert>**. The Convert Files window, Figure 8-9, is displayed. Input the drive where the file(s) are to be stored and select Lotus or ASCII format. Then click on **[OK]** to convert the selected file(s) and remove the Convert Files window from the display.

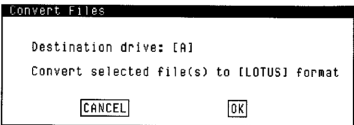


Figure 8-9. Convert Files Window

File Utilities

8.10 Receive File Command

To receive a file from an external computer through the Communications port and place it in the directory that is currently selected:

- Display the File Utilities window by clicking on "File Utilities" from the Main window.
- 2. Click on the desired directory, listed on the left-hand side of the window.
- 3. Click on <Receive>. The Receive File window, Figure 8-10, is displayed. Input the directory, file name and the extension for the incoming file. The selections for the extensions are shown in Table 8-3.

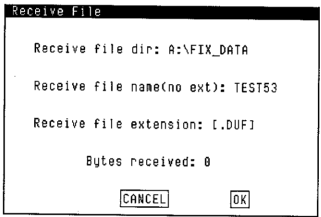


Figure 8-10. Receive File Window

Extension	File Type
.SCN	Wavelength scan data
.APX	Method or other text file
.DUF	Data file, consisting of data at selected wavelengths and calculated information.
.PGM	Program
.WKS	Lotus
.ASC	ASCII, converted from a .DUF file.
.FFT	Fourier transform data from
	Multicomponent Analysis mode.

Table 8-3. File Extensions

- 4. Click on [OK]. The cursor arrow will change to the hour glass and [CANCEL] on the bottom of the window will change to [STOP RECEPTION].
- 5. Ready the external computer to send the file. Send the data to the instrument. As the data are received by the instrument, "Bytes received" on the Receive File window is updated.
- 6. When all the data have been sent to the instrument, click on [STOP RECEPTION] to remove the Receive File window.

8.11 Transmit File Command

The Transmit command is used to transmit a file to an external computer through the Communications port. This command is also used to print a file on the Dot Matrix Printer. Before transmitting files other than Method files to the Printer, do an ASCII conversion. Only Method files and ASCII-converted files can be printed.

- Display the File Utilities window by clicking on "File Utilities" from the Main window.
- 2. Click on the desired directory, listed on the left-hand side of the window. All the files in the selected directory are listed on the right-hand side of the window.
- 3. Click on one or more file names to highlight them.
- 4. Click on <Transmit>. The Transmit File window, Figure 8-11, is displayed. Select either [Com/RS232 Port] or [Printer].



Figure 8-11. Transmit File(s) Window

- 5. Ready the external device to receive the file(s).
- 6. Click on [OK] to send the files. As each file is sent, the highlight is removed from the file listing. When all the files have been sent, the Transmit File window is removed.

SECTION NINE

DATA I/O

The Communications port is used to connect the instrument to an external computer. There are three uses for the communications port.

- 1. Output Mode To output data as they are collected. The type of data output is selected in the Communications Configuration window. All collected data are output, as long as the output parameter is enabled. Instructions are provided in section 9.3.
- 2. **Remote Control** To control the instrument remotely. Instructions are provided in sections 9.4 to 9.6.
- 3. File Transfer To receive and transmit files that are stored in the instrument. Instructions are provided in sections 8.9 and 8.10, respectively.

a

9.1 Communications Configuration

The Communications Configuration window is used to set the communications parameters for the Communications port, to enable the Output mode, and to select the type of data output for all analysis modes of the instrument. This window is also used to select the parameters for the X-Y Plotter. Instructions for setting up the Communications Configuration window for use with the X-Y Plotter are provided in Manual 514523.

To display the Communications Configuration window, Figure 9-1, enter the Configuration mode by clicking on "Configuration" from the Main window to display the Configuration window, then click on <Communications> from the Configuration window.

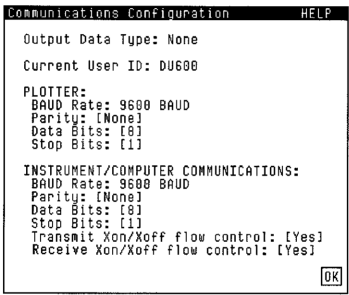


Figure 9-1. Communications Configuration Window

Input the following parameters:

1. Output Data Type - This is used to enable the Output mode. Click on to select [None], which is no output, or [Raw and calculated], [Raw] and [Calculated], which is the type of data sent to the Communications port. The format of the output data is described at the end section in this manual that describes the analysis mode.

- 2. Current User ID Input an identification, which is transmitted in the Output mode at the beginning of each data set.
- 3. Instrument Communications Parameters Set the following parameters, so that they are consistent with the remote device.
 - a. Baud Rate Click on this parameter to display a window with the selections for baud rate. Click on the desired baud rate to select it, then click on [OK] to remove the window.
 - b. Parity Toggle between [None], [Even], and [Odd] to set the parity.
 - c. Data bits Toggle between [7] and [8] to set the number of data bits.
 - d. Stop bit Toggle between [1] and [2] to set the number of stop bits.
 - e. Transmit Xon/Xoff flow control Toggle between [Yes] and [No] to enable or disable transmission of Xon/Xoff.
 - f. Receive Xon/Xoff flow control Toggle between [Yes] and [No] to enable or disable reception of Xon/Xoff.

NOTICE

When 7-bit data is being received and transmitted by the instrument, both flow control fields should be set to [Yes]. When 8-bit data is being received by the instrument, set "Transmit Xon/Xoff" to [Yes] and "Receive Xon\Xoff" to [No]. When 8-bit data is being transmitted by the instrument, set "Transmit Xon/Xoff" to [No] and "Receive Xon\Xoff" to [Yes]. Examples of 8-bit data include files with the following extensions: .WKS, .DUF, .SCN, .FFT, and .PGM.

When all the desired parameters are input, click on [OK] to store the entries and remove the Communications Configuration window from the display.

9.2 Diagnostics

Start Up Instructions

Use the following check list to verify that the Communications port is operational.

- 1. Verify that both the instrument and remote device have been powered up.
- Verify that the remote device is configured as DTE Data Terminal Equipment. (The DU-600 Spectrophotometer is configured as DCE -Data Communications Equipment.)
- 3. Select the communications parameters on the instrument (baud rate, parity, data bits, etc.), using the Communication Configuration window.
- 4. Configure the external device for the baud rate, number of data bits, number of stop bits, and the parity to correspond with the instrument.
- 5. Verify that the Communications port on the instrument is operational by performing the Loop Back Test.
- 6. Connect the external device to the Communications port using a cable. The COM port has a standard DB-25 male connector. A cable is not provided. However, one is available from Beckman, which has a standard DB-25 female connector for the DU Series 600 Spectrophotometer and a standard DB-25 male connector for the external device.

The following table shows the pin configuration of the DB-25 connector.

Pin Number	Circuit Description
1	Protective ground
2	Receive data
3	Transmit data
7	Signal ground

7. If the external device has output capabilities, verify that the instrument receives information sent to it by performing the Input Test.

Loop Back Test

To perform the loop back test on the Communications port:

1. Locate the Communications port on the back of the instrument. Disconnect the cable that is connected to it. See Figure 9-2. Insert the RS-232 Test Plug into the desired port. The RS-232 Test Plug is provided in the shipping kit with the instrument.

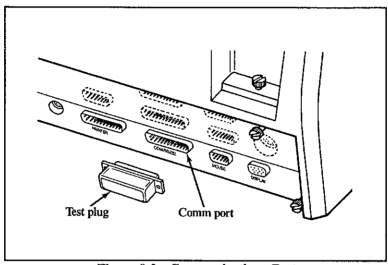


Figure 9-2. Communications Port

2. With the Main window displayed, click on "DIAGNOSTICS" to display the Diagnostics window. Then click on <LoopBack>, to display the Loop Back Tests window, Figure 9-3.

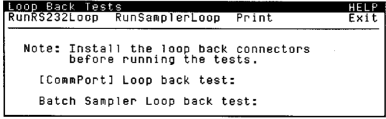


Figure 9-3. Loop Back Tests Window

Data I/O

- 3. If the desired RS-232 port is not displayed in brackets on the Loop Back Tests window, click on the bracketed selection to display the desired port, then click on [OK] to select the port and remove the window from the display.
- 4. Click on <RunRS232Loop>. The instrument automatically performs the test and displays the result, passed or failed, on the Loop Back Tests window.

NOTICE

If the test fails, the instrument is malfunctioning. Contact the local Beckman service office.

5. When the test is complete, click on **Exit** to return to the Diagnostics window.

Input Test

The RS-232 Input test verifies that the instrument is receiving the information sent by an external computer. The input test should be performed after the Loop Back test has verified that the Communications port on the instrument is operational. To perform the Input test:

- 1. Verify that the instrument and the external device have the same communication parameter settings (baud rate, parity, data bits, stop bits and Xon/Xoff). The Communications Configuration window is used to set the parameters on the instrument.
- 2. Locate the Communications port on the back of the instrument. Connect the external device to it.
- 3. With the Main window displayed, click on "DIAGNOSTICS" to display the Diagnostics window. Then click on <RS232Input> to display the RS232 Communications Input window, Figure 9-4. The bottom of the window is blank.

R\$232 Commus	nications Input	;	
Print			Exit
Printable (ASCII followed	by hex value	
		_	

Figure 9-4.
RS232 Communications Input Window

4. Send an ASCII code from the external device to the DU-600 Spectrophotometer. The ASCII and corresponding hex value of the input received by the instrument are displayed.

NOTICE

If nothing is displayed, the external device is not communicating. Use the Loop Back Test to verify that the instrument Communications port is operational. If the wrong values are displayed, the communication parameters are probably different on the two devices.

5. When the test is complete, click on **Exit** to return to the Diagnostics window.

9.3 Output Mode

The Output mode is used to output data to the Communications port as they are collected by the instrument. The user can select to output only the raw data, only the calculated data, or both the raw and the calculated data. This data output selection is made on the Communications Configuration window, Figure 9-1. A description of the exact information that are sent for "raw", "calculated" and "raw and calculated" is provided near the end of each operating mode.

A header is sent prior to the data. The header consists of four lines, each followed by a carriage return and a line feed. The first line is the User ID, which is input in the Communications Configuration window. The second line is the Method ID, which is the method file which is used to collect the data. The third and fourth lines are the date and time, respectively.

9.4 Remote Control Mode

In the Remote Control mode, all instrument functions are controlled by a remote device through the Communications port.

NOTICE

Angle brackets "<" and ">" are used throughout this section to indicate the start and end of a remote command, respectively. These brackets are not part of the command and should not be sent with the command.

To operate the remote control mode:

- 1. Verify that the instrument and the external device are communicating. Refer to section 9.2 for additional information.
- With the Main window displayed, click on "REMOTE CONTROL" to enter the Remote Control mode. A window with the message "Remote control operation, click on any button to quit" is displayed. No additional instrument function is required.

To enter the Remote Control mode from the remote device, with either the Power Up Diagnostics window or the Main window displayed, the remote device must send the following two ASCII codes to the instrument:

<ctrl W> <ctrl V>
 or
(decimal 23) (decimal 22)

3. The instrument sends <r> <lf>, indicating that it is ready to receive remote commands from the remote device. The format for a remote command is:

<ascii string> <lf>,

where, <ascii string> is a string of up to 40 ASCII characters, and <lf> is a line feed. The remote commands are described in the next section. All non-printing commands, other than <lf>, are ignored by the instrument.

4. When the instrument receives a command, it returns <s> <lf>, if it successfully executes the command. If the command cannot be executed, <f> <lf> or <lf> is returned.

5. To exit the Remote Control mode from the instrument, click on a mouse button. A warning window is displayed. When [OK] is clicked on, the instrument sends <x> <lf>, indicating that the instrument is exiting the Remote Control mode.

To exit the Remote Control mode from the external device, send the <exit> command.

NOTICE < lf> is < ctrl J>.

9.5 Remote Control Commands

The following are the remote commands that the instrument can recognize. Each command must be followed with a <|f>.

NOTICE

The remote commands frequently use values stored in the x-, y-, z- and t-registers in the stack. For simplicity, values stored in these registers are referred to as x-, y-, z- and t-values. That is, the x-value is the value stored in the x-register.

Math Commands

Command	Description
<number></number>	Any valid number. It is placed in the x-register.
<enter></enter>	Push the stack. X-value goes to y-register, y-value goes to z-register, z-value goes to t-register, t-value is lost. Original x-value remains in x-register, and is overwritten when subsequent number is input.
<rotate></rotate>	Rotate all the values in the stack: $t\rightarrow z$, $z\rightarrow y$, $y\rightarrow x$, $x\rightarrow t$.
	In/Out Commands
<rs232_out></rs232_out>	Send the x-value to the Communications port.
<rsbyte_out></rsbyte_out>	Convert the x-value to a single byte and send to the Communications port.

Spectro Commands

Command	Description
<source/>	Turn on or off the desired source, determined by the x-value. If: x = 0, visible off, x = 1, visible on, x = 2, UV off, and x = 3, UV on.
<set_wl></set_wl>	Set the wavelength to the x-value.
<set_rd_avg></set_rd_avg>	Set the read average to the x-value.
<black></black>	Blank at the wavelengths given in registers 30 to 41. The x-value determines the total number of wavelengths.
<read_abs></read_abs>	Take an absorbance reading using the <set_wl> and <set_rd_avg> values. Place the reading in the x-register. (Blank the instrument at the desired wavelength before taking a reading.)</set_rd_avg></set_wl>
<read_%t></read_%t>	Take a transmittance reading using the <set_wl> and <set_rd_avg> values. Place the reading in the x-register: (Blank the instrument at the desired wavelength before taking a reading.)</set_rd_avg></set_wl>
<set_wl_range></set_wl_range>	Set the wavelength range for scanning, where the lower wavelength is placed in the y-register and the upper wavelength is placed in the x-register.
<set_spd></set_spd>	Set the wavelength scanning speed, determined by the x-value. If: x = 0, the speed is 120 nm/minute, x = 1, the speed is 240 nm/minute, x = 2, the speed is 600 nm/minute, x = 3, the speed is 1200 nm/minute, and x = 4, the speed is 2400 nm/minute.

Perform a blank scan over the designated wavelength

range at the designated speed.

<scan_blank>

Command	Description
<scan_abs></scan_abs>	Perform an absorbance scan over the designated wavelength range at the designated speed. Place the readings in the designated buffer locations.
<scan_%t></scan_%t>	Perform a transmittance scan over the designated wavelength range at the designated speed. Place the readings in the designated buffer locations.
<send_scan></send_scan>	Send the contents of the buffer ¹ to the remote device. The wavelength range is determined by <set_wl_range>.</set_wl_range>
<put_buffer></put_buffer>	Place the x-value in the buffer at the location determined by the y-value.
<get_buffer></get_buffer>	Find the value in the buffer location determined by the x-value and place it in the x-register.
<recv_file></recv_file>	The file must be a Method file or a data file previously converted to an ASCII file. The command must be followed by a <space>, then the complete pathname of the file. The instrument will receive the file and store it in the designated location. The instrument expects to receive an EOF (hex 1a) to indicate the end of the file.</space>
<wl_calib></wl_calib>	Perform a wavelength calibration.

¹ The readings from a wavelength scan are stored in buffer locations 1900 to 11000, which when multiplied by 0.1 represent the wavelength. At a scanning speed of 120 nm/minute, each buffer location is filled. At faster speeds, not all of the buffer locations will be filled. The locations that are filled are shown in the following table.

Scan Speed	Delta 1/Reading	Delta Buffer Locations
2400	2	20
1200	1	10
600	0.5	5
240	0.2	2
120	0.1	1

For example, the first five readings of a scan taken from 400 to 500 nm at 240 nm/minute are stored in buffer locations 5000, 4998, 4996, 4994, 4992.

Command

Description

<send file>

The file must be a Method file or a data file previously converted to an ASCII file. The command must be followed by a <space>, then the complete pathname of the file. The instrument will send the designated file. The instrument will send an EOF (hex 1a) to indicate the end of the file.

<beep>

Make an audible beep.

<exit>

Exit the Remote Control mode.

<ex_mode>

Exit the Remote Control mode and display the first window of the analysis mode that is indicated by the mnemonic following this command. The analysis mode will operate in the normal manner. When operation of the analysis mode is complete and <Quit> is clicked on, the instrument will return to the Remote Control mode.

Mnemonic Analysis Mode

fix	Fixed Wavelength
	Wavelength Scan ²
kin	Kinetics/Time
prot	Protein Analysis
frac	Fraction Read/Plot
na	Nucleic Acid
sca	Single Component Analysis
em	Enzyme Mechanisms
ea	Enzyme Activity
mca	Multicomponent Analysis
gela	Gel Scan Area
gelm	Gel Scan Molecular Weights
pv	Performance Validation ³
diag	Diagnostics

²The Wavelength Scan mode cannot be entered from the Remote Control mode.

³In this mode only, the tests are performed automatically, the results are output to the printer, and the Remote Control mode is re-entered. About 75 minutes are required to complete the Performance Validation tests.

Command Description

<set_output> When using <ex_mode> to operate an analysis mode, this selects the output mode, determined by the x-value. If:

x = 0, no output,

x = 1, raw and calculated output,

x = 2, raw data output, and

x = 3, calculated data output.

<pwrup_res>

Send the instrument serial number and results of the Power Up Diagnostics to the remote device.

Register Commands

<store_reg> <#> Store the x-value in the register designated by the
number following this command.

Accessory Commands

<sip_fill>

Run the sipper pump in the forward direction for the time, in seconds, indicated in the x-register at the speed indicated in the y-register. (The speed range is 1 to 10, where 10 is fastest.) If the x-value is 0, the fill time input in the Accessory Configuration window is used. If the y-value is 0, then a speed of 5 is used.

<sip_return>

Run the sipper pump in the reverse direction for the time, in seconds, indicated in the x-register at the speed indicated in the y-register. (The speed range is 1 to 10, where 10 is fastest.) If the x-value is 0, the return time input in the Accessory Configuration window is used. If the y-value is 0, the speed is 5.

<adv_batch>

Advance the batch sampler one tube position.

<raise_arm>

Raise the batch sampler aspirator arm.

Command Description <lower arm> Lower the batch sampler aspirator arm. <to wash> Move the batch sampler aspirator arm to the wash station. <to samp> Move the batch sampler aspirator arm from the wash station back to the sample position. <get temp> Read the current temperature and place it in the xregister. <set temp> Turn on the temperature controller and set the temperature to the value currently in the x-register. If the value is outside the range of 10.0 to 90.0 (or 0.0 to 110.0 for the Tm cell holder), the temperature controller turns off. Move the transport to the millimeter position currently <set mm> in the x-register. Move the transport to the cell position currently in the <set cell> x-register. To home up the transport, place 13 in the x-register. <sampler> 8-channel Pump - Control is based upon the x- and yvalues, where: x = 0 to stop the pump or 1 to start the pump y = determines pump direction, where 0 is reverse

v-values, where:

(return) and 1 is forward (fill). 16-position Valve - Control is based upon the x- and

9.6 Remote Examples

EXAMPLE 1

Set up the instrument to take fixed wavelength readings at 280 nm every 10 seconds on a flowing sample and send the data to the remote device.

SOLUTION

The remote device must do the timing and request the readings. The following are the remote commands:

Command	Description
<1> <lf></lf>	Set the x-register to 1.
<set_rd_avg><lf></lf></set_rd_avg>	Set the read average to 1 second.
<280> <lf></lf>	Set the x-register to 280.
$<$ set_wl $>$ <lf<math>></lf<math>	Set the wavelength to 280.

Start blank solution flowing through the flowcell.

Start the sample flowing through the flowcell.

Repeat the last two commands every 10 seconds to take readings on the flowing sample.

EXAMPLE 2

Set up the instrument to take fixed wavelength readings at 260 and 280 nm on samples placed in the Auto 6 Cell Holder. Send the readings to the remote device.

SOLUTION

The following are the remote commands:

Description
Set the x-register to 260.
Store 260 in register 30.
Set the x-register to 280.
Store 280 in register 31.
Set the x-register to 1.
Set the read average to 1 second.

Place blank solution in cell position 1.

<1> <lf></lf>	Set the x-register to 1.
<set_cell><lf></lf></set_cell>	Move to cell position 1.
<2><1f>	Set the x-register to 2.
<black><lf></lf></black>	Take a blank reading at the two wavelengths.

Place the first six samples in the Auto Cell Holder.

<260><1f>	Set the x-register to 260.
<set_wl><lf></lf></set_wl>	Set the wavelength to 260.
<read_abs><lf></lf></read_abs>	Take a sample reading.
<rs232_out><lf></lf></rs232_out>	Send the sample reading to the remote device.
<280><1f>	Set the x-register to 280.
<set_wl><lf></lf></set_wl>	Set the wavelength to 280.
<read_abs><lf></lf></read_abs>	Take a sample reading.
<rs232_out><lf></lf></rs232_out>	Send the sample reading to the remote device.
<2><1f>	Set the x-register to 2.
<set_cell><lf></lf></set_cell>	Move to cell position 2.

Repeat the above 10 steps, each time incrementing the <set_cell> command to move the transport.

EXAMPLE 3

Using a batch sampler, blank on the first tube and make a wavelength scan from 350 to 700 nm at 1200 nm/minute on the next three tubes. Send the data to the remote device.

SOLUTION

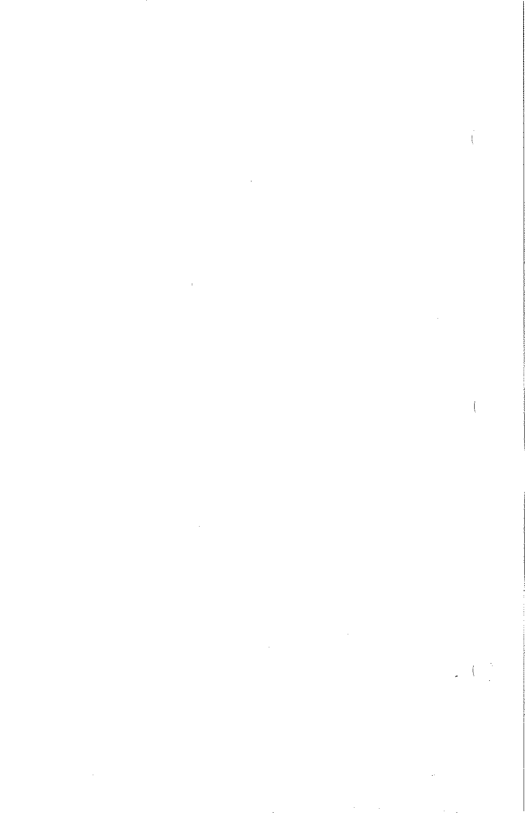
Command	Description
<350> <lf></lf>	Place 350 in the x-register.
<enter><lf></lf></enter>	Move it into the y-register.
<700><1f>	Place 700 in the x-register.
<set_wl_range><lf></lf></set_wl_range>	Set the wavelength scanning range from 350 to 700 nm.
<1><1f>	Place 1 in the x-register.
<set_spd><lf></lf></set_spd>	Set the scan speed to 1200 nm/minute.

These steps are for the blank.

<lower_arm> <lf></lf></lower_arm>	Lower the aspirator arm into the blank solution.
<8> <lf></lf>	Input 8 in the x-register.
<enter><lf></lf></enter>	Move it to the y-register.
<5> <lf></lf>	Input 5 in the x-register.
<sip_fill><lf></lf></sip_fill>	Aspirate the blank for 5 seconds at a pump speed of 8.
<scan_blank><lf></lf></scan_blank>	Scan the blank.
<raise_arm><lf></lf></raise_arm>	Raise the aspirator arm.
<sip_return><lf></lf></sip_return>	Return the blank using the same time and speed as the fill.

Repeat the following steps for each sample.

<adv_batch><lf></lf></adv_batch>	Advance to the next tube.
<lower_arm><lf></lf></lower_arm>	Lower the aspirator arm into the first sample.
<sip_fill><lf></lf></sip_fill>	Aspirate the first sample for 5 seconds at a pump speed of 8 (the values are still in the
	registers).
<scan_abs><lf></lf></scan_abs>	Scan the first sample.
<send_scan><lf></lf></send_scan>	Send the scan for the first sample to the remote device.
<raise_arm><lf></lf></raise_arm>	Raise the aspirator arm.
<sip_return><lf></lf></sip_return>	Return the sample.



PREVENTATIVE MAINTENANCE

10.1 General Information

The DU Series 600 Spectrophotometer is a microprocessor controlled, optical instrument. For the microprocessor to operate properly, it is necessary to have adequate ventilation, as described in the installation instructions for the instrument, Manual 517315. It is necessary to operate the instrument below the specified maximum temperature of 40°C (104°F) for adequate cooling.

For the optical components to retain their quality, it is necessary to operate the instrument in a dust-free, non-corrosive environment. There can be situations where a spectrophotometer must be placed in a dust or corrosive environment. In this case, the optical surfaces must be cleaned or replaced periodically. Contact Beckman service for more information.

One component of the DU Series 600 Spectrophotometer that can be user-serviced to prevent an interruption in operation or increase performance is the UV source. With time, the output of the UV source gradually decreases. Typically, the output decreases to one-half the original level in 1000 hours of use. Decreased output from the UV source decreases the sensitivity of the instrument. This effect is wavelength dependent. If the sensitivity of the instrument decreases below the necessary level and the source has been on for 1000 or more hours, the UV source should be replaced, even if it is not burned out. The length of time that the UV source has been illuminated is displayed by clicking on <Status> from the Diagnostics window.

If the performance of the instrument remains satisfactory, the UV source does not need to be changed until it burns out. A burned out source is indicated by the "UV LAMP Failure" message.

Decreased output from the visible source is normally not significant enough to cause decreased sensitivity. However, if decreased sensitivity is observed in the visible range, the visible source should be checked for a brown film coating inside the lamp. To check the source, remove it using the instructions in section 12.3. If the lamp has a brown coating, replace it with

a new lamp. If the lamp is clear, it is operational. Replace the source, then peak it using the instructions in part II of section 12.3.

A service contract for this instrument is available from the Beckman field service representative.

10.2 Status Window

The Instrument Status window is used to display the cumulative total amount of time that the instrument has been turned on, the cumulative total amount of time that the UV source has been on (the time for the UV source can be reset in the Diagnostics mode when the source is replaced), the serial number of the instrument and the software revision.

To display the Instrument Status window:

- 1. With the Main window displayed, click on "DIAGNOSTICS" to display the Diagnostics window.
- 2. Click on **<Status>**. An Instrument Status window, similar to the one in Figure 10-1, is displayed.

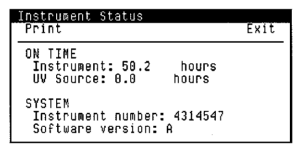


Figure 10-1. Status Window

- 3. When the desired information is obtained, click on **Exit>** to remove the Instrument Status window.
- Click on <Quit> to remove the Diagnostics window and display the Main window.

TROUBLESHOOTING

11.1 Power Up Diagnostics

Each time the instrument is powered up, a series of diagnostic tests is performed automatically to ensure proper operation of major system components. As the tests are performed, a window with "Executing Power Up Diagnostics" is displayed. When all tests are complete, the Power Up Diagnostics window is displayed.

Each of the diagnostic tests is listed on the Power Up Diagnostics window. Following the test name is "Passed", "Failed" or "Not Installed". If any of the tests fail, power down the instrument, then power it up again. If the failure persists, the instrument may be partially operational, as described in the following table.

The "Visible Lamp" and "Light Path" failures can be corrected by the user. (If the problem persists, contact the local Beckman service office.) If any other test fails, contact the local Beckman service office.

Failed Test

Explanation

CPU
PROM
RAM Controller
RAM
Video Controller
Video RAM
Video Palette

RS232 Ports 1 and 2

The instrument cannot be used.

RS-232 ports 1 (mouse) and/or 2 cannot be used.

EE PROM

Configuration parameters and the wavelength calibration will not be stored when the instrument is powered down. Calibrate the wavelength each time the instrument is powered up if this test fails.

Failed Test

Explanation

PROM Option Software Option

Software options are not available.

RAM Option

The expanded memory option in the A-drive is not available.

RAM Battery Backup

The files stored in the A-drive will be lost when the power is turned off.

Programmability

Programmability is not available.

RS232 Ports 3 and 4

RS-232 ports 3 and/or 4 cannot be used.

Keyboard Processor

The keyboard cannot be used.

Detector Gain Readings cannot be taken. The instrument can be used for other functions.

Visible Lamp

Replace the visible source. Directions are provided in section 12.3 of this manual.

Light Path

Something is blocking the light path through the sample compartment. Remove the sampling accessory and power up the instrument again.

Filter Lamp Selector Readings cannot be taken. The instrument can be used for other functions.

Wavelength Drive

The wavelength may not be accurate. Use the instrument only if the accuracy of the readings can be verified. Contact Beckman service.

System Clock

The date and time display may not be accurate. Also, time intervals may not be timed correctly.

11.2 Operational Failures

There are three failures that can occur during instrument operation. These are displayed under the Permanent Menu Bar on the left-hand side. The failure is displayed until the condition is corrected.

Message	Recommended Action
VIS LAMP Failure	Replace the visible source. Directions are provided in section 12.3 of this manual.
UV LAMP Failure	Replace the UV source. Directions are provided in section 12.2 of this manual.
OVER TEMPERA- TURE SHUTDOWN	Contact the local Beckman service office. Do not continue using the instrument.

11.3 Operational Messages

The following messages can occur while operating the instrument and are displayed in an error window. In most cases, these errors do not indicate an instrument malfunction, but a situation where an action was initiated that the instrument cannot perform.

If the instrument can continue operating with the error condition, the window is removed after 3 seconds. If the instrument requires operator action to continue, the user must click to remove the window.

Message	Explanation/Recommended Action
Bad file name	The input file name is invalid. Check the characters input for both the directory and file name.
Cannot execute custom application	The method directory where the application was originally created is full. Delete files from the method directory to make more room.
Could not write to disk	Check the disk. It is not installed, is full, or is write protected.
Cannot write to temporary file	Data cannot be stored using a default file name. Input a different file name.
Changes not saved, File protected	The method file is protected. To store the changes, protection must be removed.
Delete failed	The file to be deleted could not be found; verify the disk is installed in the drive. Protected method files cannot be deleted.
Disk controller fault	Check the disk drive. It is not installed or no disk is in the drive.
Drive error	Check the disk drive. It is not installed or no disk is in the drive.
Drive timeout	There is no disk in the drive. Insert a disk in the drive.

Message Explanation/Recommended Action End of file The file has an error in it. No data exists at the location specified by the row and column. Exceeded number of The maximum number of readings that readings can be stored in a file has been reached. Click on <SaveClear> to either store or delete the data. Then data collection can resume. Exceeded number of The number of annotations that can be notes (4) made on a graph is limited to 4. Failed to save method There is not enough room on the disk to file store the file. Insert another disk or delete file(s) to make room on the current disk. File already exists The user asked for data to be stored in a file that already exists. If the data are stored, the existing data will be overwritten. Or, input a different file name. File is protected If the method file is protected, the file cannot be deleted and changes to the file cannot be stored. Remove protection to change the file.

File not found

Either the source or destination file was not found. If disk was removed or exchanged after the directory was displayed, replace the original disk.

File not saved

The method/data were not stored. Check the disk. There is no space on the disk, a faulty disk, or the file name is invalid.

File protection was not changed

An unacceptable user name or password was input. Input the correct user name or password.

Message

Explanation/Recommended Action

File Read Failed

The data were not retrieved. Check the disk. There is no disk, no space on the disk, a faulty disk, or the file was not closed properly.

File Write Failed

The data were not stored. Check the disk. There is no disk, no space on the disk, a faulty disk, or the file is already closed.

The disk was not formatted. Check the disk. It may be faulty or the density may be wrong.

Illegal file name

Format failed

A default file name cannot be used for a data file. Reserved file names include: "WORK RES", "WORK STD", "WORK", W1 RES", and "SCAN". Use another file name.

Incompatible file type

The file cannot be read because it was copied into a directory other than where it was created.

Invalid directory

The file cannot be moved/copied into the selected directory.

Invalid disk

Check the disk. It is not formatted or is faulty.

Memory allocation failed

On entry to an analysis mode, the system was unable to find enough memory to Reset the instrument. This can result from a very long program executing an analysis mode.

Method directory full

A maximum of 100 method files can be stored in a directory. Delete existing files to make room for additional files.

Move failed There was not enough room to store the file in the new location. Delete file(s) to make more space. No disk controller The disk drive is not installed. Out of disk space The disk is full. Insert a new formatted dick OVER TEMPER-Contact the local Beckman service office. ATURE SHUTDOWN Do not continue using the instrument. Printer is not On Line Press the "ON LINE" button on the printer, so the indicator light is illuminated. Program area full A program can have a maximum of 1000 steps. Edit the program so that it has less steps. Read limit reached, click The maximum number of readings has on SaveClear been reached. Click on <SaveClear> to either store or delete the data. Then data collection can resume. Rename failed The file was not renamed because it could not be found or an invalid file name was input. Check the file names. Reserved file name A reserved file name cannot be used to store data. Reserved file names include: "WORK_RES", "WORK_STD", "W1 RES", "WORK", and "SCAN". Change the file name.

names.

Too many files selected for renaming

Source and Destination

files are the same

Message

More than one file name was clicked on for renaming. Only one file can be renamed at a time.

A file cannot be moved, copied or re-

named if the source and destination files are the same. Change either of the file

Explanation/Recommended Action

Message

Explanation/Recommended Action

Transport failure

The transport cannot move to the expected location. Verify that the number of cells input in the analysis parameters is not greater than the number of cells in the transport configuration. Verify that nothing is blocking the movement of the transport, then "HOME" the transport using the Device Control window. If the failure persists, contact Beckman service.

Transport not installed

The Auto Cell Holder cannot be used because the transport is not installed. If it is physically in the sample compartment, verify that the cable is connected to the back of the instrument.

Unable to convert file to ASCII format

Method files are already ASCII files and therefore do not need to be converted to ASCII files.

Unable to convert file to Lotus format

Method files cannot be converted to Lotus.

Unable to open destination file Verify the file name is valid and that the disk is installed.

Unable to open source file

Verify the file name is valid and that the disk is installed.

UV LAMP Failure

Replace the visible source. Directions are provided in section 12.3 of this manual.

VIS LAMP Failure

Replace the UV source. Directions are provided in section 12.2 of this manual.

Warning: Blank Needed

Place a cuvette of solvent in the cell holder and click on << BLANK>>. Any reading taken without a blank is invalid.

Message

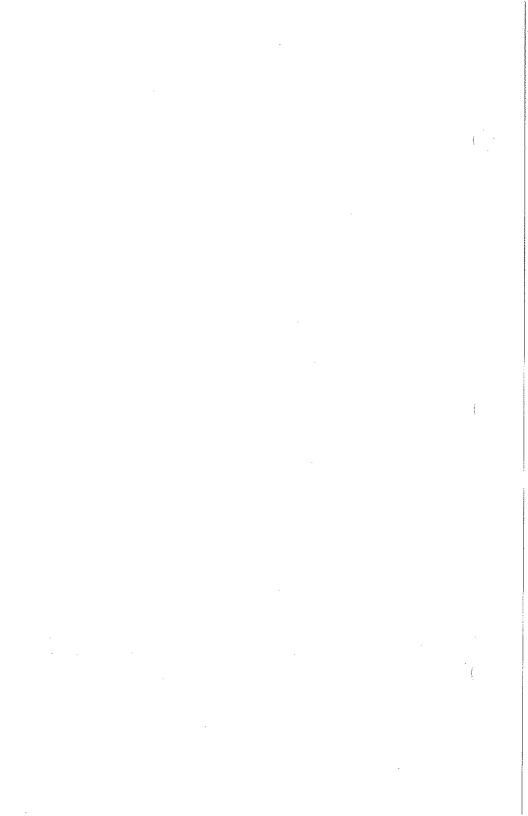
Explanation/Recommended Action

Warning: Lamps Off

Turn on the source(s). Wait for the UV source to fire, then blank the instrument. Any reading taken without a source on is invalid.

Wrong file type

The file type is incorrect for the mode being used. The file was copied into an incompatible directory.



CORRECTIVE MAINTENANCE

A NOTICE

Refer to the SAFETY NOTICE before following a procedure in this section. Refer all servicing for procedures not contained in this section to qualified Service personnel.

12.1 Fuse Replacement

The only user accessible fuses are the power fuses, located next to the receptacle where the power cord is plugged into the instrument. The instrument has one fuse if it is 100/120V; it has two fuses if it is 220/240V.

Parts required:

100/120V instrument - 3A slow blow fuse (UL/CSA), P/N 883908 220/240V instrument - 1.6A slow blow fuse (IEC), P/N 897176

Tools required: Small flatblade screwdriver

- 1. Turn the instrument around, so that the back can be accessed.
- 2. Unplug the power cord from the instrument.
- 3. Use the end of the screwdriver to pry the cover off of the fuse holder. See Figure 12-1.

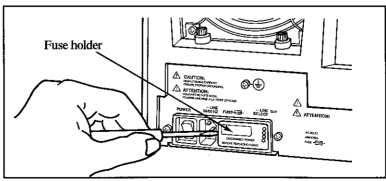


Figure 12-1.

4. Note the orientation of the fuse(s) in the holder. Remove the fuse, then replace with the new fuse, oriented in the same position. See Figure 12-2. On 220/240V instruments, both of the fuses should be replaced, even though only one may look faulty.

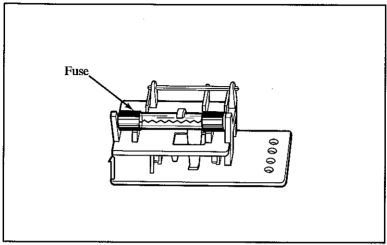


Figure 12-2.

△ WARNING

For continued protection against risk of fire, replace the fuse(s) only with the type and current rating specified above.

ATTENTION

Afin d'assurer une protection permanente contre les risques d'incendie, remplacer uniquement par un fusible de même type et valeur.

- 5. Replace the fuse holder, and press to secure.
- 6. Plug the power cord back into the instrument.
- 7. Turn the instrument around so that it is facing forward.

12.2 UV Source Replacement

Parts required: UV source, P/N 514366

1. Unplug the instrument power cord. Allow the instrument to cool for 15 minutes.

△ CAUTION

To avoid risk of electrical shock, disconnect power to instrument before changing either source.

The UV lamp operates at a high temperature. To prevent burns, allow at least 15 minutes for the instrument and sources to cool before handling internal components.

The UV lamp generates UV light. Do not look directly at an operating lamp without wearing UV protective eye glasses.

ATTENTION

Risque de choc électrique. Débrancher l'appareil avant de changer de source.

2. Remove the display from the top of the instrument cover. Unscrew the thumb screw which secures the source cover. Open the source cover and remove it. See Figure 12-3.

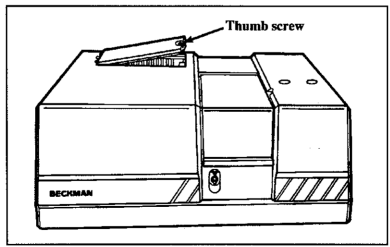


Figure 12-3.

3. To unplug the connector for the UV source, squeeze on the tabs on the sides of the connector, then pull up to release. See Figure 12-4.

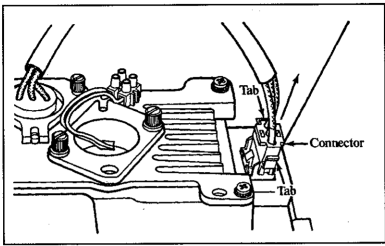


Figure 12-4.

4. Unscrew the two thumb screws which hold the UV source in position. See Figure 12-5.

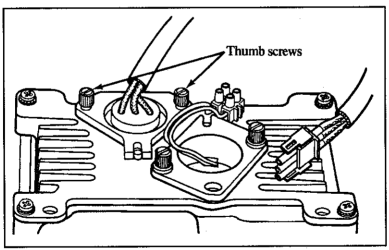


Figure 12-5.

5. Remove the source by lifting straight up. See Figure 12-6.

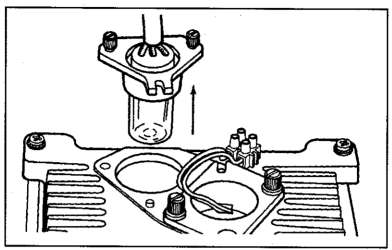


Figure 12-6.

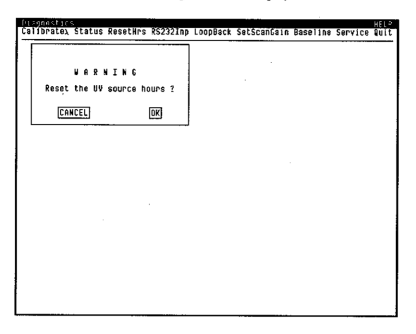
6. Unpackage the new source, being careful not to touch the glass envelope on the source. Carefully push the new source straight down, until it is seated flat against the source housing. Secure by tightening the two thumb screws, shown in Figure 12-5. Plug in the connector, shown in Figure 12-4.

NOTICE

Do not touch the glass envelope on the new source. If it is touched, clean with alcohol.

- 7. Replace the source cover and secure by tightening the thumb screw. Reposition the display on the top of the instrument cover.
- 8. Plug in the instrument. When the diagnostic checks are complete, the Power Up Diagnostics window is displayed. Click on <Quit> to display the Main window.

9. Click on **<Diagnostics>** to display the Diagnostics window. Click on **<ResetHrs>**. The following window is displayed.



10. Click on [OK] to reset the UV source hours to 0, then click on <Quit> to return to the Main window.

12.3 Visible Source Replacement

Part I. Replacing the Source

Parts required: Visible source, P/N 945672

 Unplug the instrument power cord. Allow the instrument to cool for 15 minutes.

△ CAUTION

To avoid risk of electrical shock, disconnect power to instrument before changing either source.

The visible lamp operates at a high temperature. To prevent burns, allow at least 15 minutes for the instrument and sources to cool before handling internal components.

ATTENTION

Risque de choc électrique. Débrancher l'appareil avant de changer de source.

2. Remove the display from the top of the instrument cover. Unscrew the thumb screw which secures the source cover. Open the source cover and remove it. See Figure 12-7.

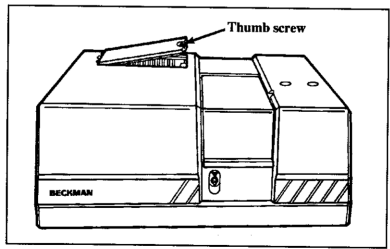


Figure 12-7.

3. Unscrew the two thumb screws which hold the visible source mounting bracket in position. See Figure 12-8.

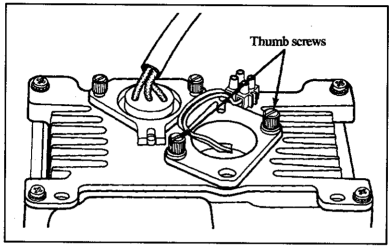


Figure 12-8.

4. Remove the source mounting bracket by lifting straight up. Rotate the source mounting bracket so that the visible source is accessible. See Figure 12-9.

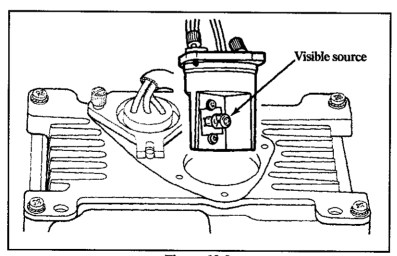


Figure 12-9.

5. Carefully remove the old source by pulling straight out. Unpackage the new source, being careful not to touch the glass envelope on the source. Hold the source with a tissue or Kim-Wipe and insert it into the source mount in the same location as the old source. See Figure 12-10.

NOTICE

Do not touch the glass envelope on the new source. If it is touched, clean with alcohol.

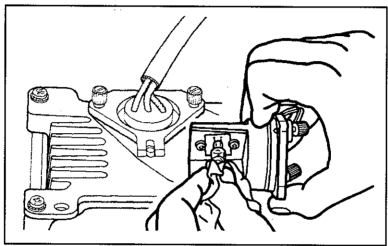


Figure 12-10.

6. Replace the source mounting bracket and tighten the two thumb screws. See Figure 12-8.

art II. Peaking the Source

Tool required: Peaking tool (Provided in the shipping kit.)

- 1. Plug in the instrument.
- 2. Click on **Exit**> to remove the Power Up Diagnostics window from the display.
- 3. Click on <<VIS OFF>> to turn on the visible source.
- 4. Click on "FIXED WAVELENGTH" to display the Fixed Wavelength window.
- 5. Click on <Parameters>, located in the menu bar at the top, center of the Fixed Wavelength Window. The Parameters window, Figure 12-11, is displayed.

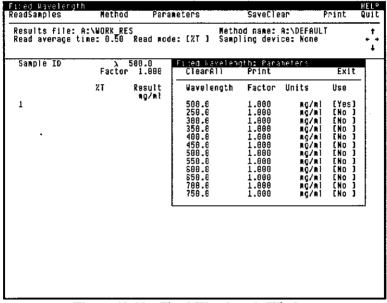


Figure 12-11. Fixed Wavelength Window with Parameters Window

- 6. To select a wavelength of 500 nm:
 - a. Click on the first wavelength value in the table and input a wavelength of 500 nm.
 - b. In the "Use" column, set the selection for the first wavelength (500 nm) to [Yes]. Set the selection for all the other wavelengths to [No].
 - Click on <Exit> to remove the Parameters window.
- 7. Locate the "Read mode" parameter, near the top in the center of the window. Verify that [%T] is displayed. If [Abs] is displayed, click on it to display [%T].
- 8. Click on < ReadBlank> to blank the instrument. A reading of 100 %T is displayed with the status information in the Permanent Menu Bar.
- 9. Loosen slightly the two thumb screws that secure the visible source mounting bracket.
- 10. Insert the peaking tool into the adjustment hole, on the front corner of the source mounting bracket. See Figure 12-12.

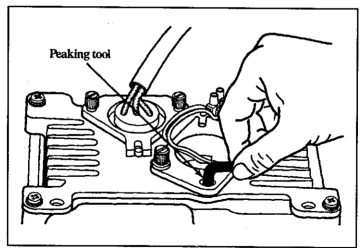


Figure 12-12.

- 11. Turn the peaking tool a small amount in the clockwise direction and observe the reading at the bottom of the display.
 - a. If the reading is higher than the initial reading, continue turning clockwise and taking readings until the maximum reading is obtained.
 - b. If the reading is lower than the initial reading, turn the tool counter-clockwise and take readings until the maximum reading is obtained.
 - c. If the reading exceeds 150%T, click on < ReadBlank> to reset the reading to 100%, then continue until the maximum reading is obtained.
- 12. Tighten the two thumb screws that secure the visible source mounting bracket. See Figure 12-8.
- 13. Replace the source cover and secure by tightening the thumb screw. Reposition the display on the top of the instrument cover.

13

SECTION THIRTEEN

TECHNICAL SPECIFICATIONS

13.1 Performance Specifications

Scan Speeds

120, 240, 600, 1200, 2400 nm/min

Data Collection Rate

20 samplings/second

Response Time

0.05 second

Wavelength Range

190 to 1100 nm

Wavelength Accuracy at 656.1 nm Full Range

±0.2 nm

Wavelength Repeatability at 656.1 nm Full Range

 $\pm 0.1 \text{ nm}$ $\pm 0.2 \text{ nm}$

 $\pm 0.5 \text{ nm}$

Spectral Bandwidth (from 200 to 680 nm)

≤1.8 nm

Photometric Readout

-0.300 to 3.000 A or 0.0 to 200.0 %T

Photometric Accuracy (at 1 A with NIST 930D solid filter at 546 nm) ±0.005 A

RMS Noise

≤0.0002 A rms¹/ ≤0.0005 A rms

(0A, average of 10 standard deviations of 10 readings at 0.05 sec intervals at 500 nm)

¹ Instruments with serial numbers 4319100 and higher.

Stray Light
(measured using NaI at
220 nm per ASTM E387-84)

< 0.05%

Stability (0A, constant ambient conditions, measured for 1 hour at 340 nm)

<0.003 A

Baseline Flatness

 $\pm 0.001 \text{ A rms}^2/$ $\pm 0.003 \text{ A rms}$

(from 200 to 900 nm, at 0 A)

13.2 Physical and Environmental Specifications

Width 69 cm (27 inches)

Height 58 cm (23 inches)

Depth 53 cm (21 inches)

Weight 33 kg (73 lbs)

Line Voltage 100/120V±10% or 220/240V±10%

Frequency 50/60 Hz

Power 200 watts typical

Ambient Temperature
Operating Range +15 to 40°C (59 to 104°F)

Humidity <85% maximum relative humidity, not to exceed 32.5°C WBT

² Instruments with serial numbers 4319100 and higher.

13.3 Storage and Transport

The DU Series 600 Spectrophotometer should be stored with the power off when not in use for several days or more. The sample compartment cover should be closed. It is necessary for the instrument to be placed in the upright position on a flat surface, or the optics could become misaligned.

The instrument should be transported in the upright position, with care taken not to jolt, bounce or shake the instrument in transport. Pack the instrument in the original shipping container for transport, if available.

13.4 Sample Compartment Configuration

The diagram in Figure 13-1 shows the location and size of the beam in the sample compartment. Notice that the beam focus is near the right-hand side of the sample compartment. Sampling accessories, which are installed correctly, will position the sample at the beam focus.

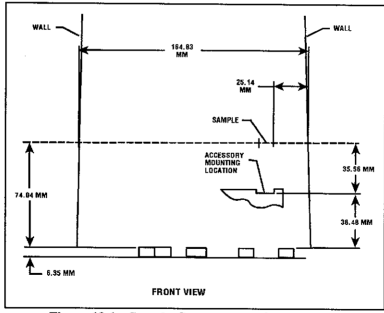
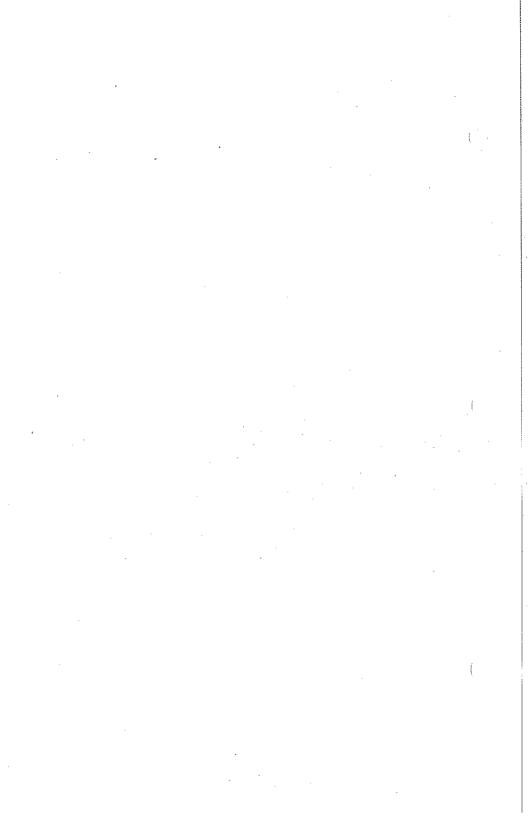


Figure 13-1. Sample Compartment Configuration

Specifications 13-3



SECTION FOURTEEN

PARTS, SUPPLIES AND ACCESSORIES

		
Instruments		
DU-640 Spectrophotometer, monochrome, 100/120V	517000	
DU-640 Spectrophotometer, monochrome, 220/240V	517004	
DU-640 Spectrophotometer, color, 100/120V	517002	
DU-640 Spectrophotometer, color, 220/240V	517006	
DU-650 Spectrophotometer, monochrome, 100/120V	517001	
DU-650 Spectrophotometer, monochrome, 220/240V	517005	
DU-650 Spectrophotometer, color, 100/120V	517003	
DU-650 Spectrophotometer, color, 220/240V	517007	
System Accessories		
Printer Options		
Dot Matrix Printer	523400	
X-Y Plotter, 115V	523413	
X-Y Plotter, 230V	523414	
System Options		
Keyboard	514666	
Bar Code Accessory	514243	
3½" Disk Drive, Controller Board and Cable	514244	
1 MEG RAM	514252	
Software Option Board	517035	
Software Options		
Protein Assay	517025	
Nucleic Acid Analysis	517026	
Fraction Read/Plot	517027	
Performance Validation	517029	
Single Component Analysis	517030	
Multicomponent Analysis	517031	
Enzyme Mechanism	517032	
Enzyme Activity	517033	
Gel Scan	517034	

Sampling Accessories

Accessory Option Board	517036
Sample Holder Mount Options Stationary Mount Transport	517023 514241
Sipper Options Sipper Accessory Unheated Flowcell Accessory Adjustable Cell Holder Micro Sipper Accessory	514242 514253 594439 514256
Batch Sampler Options Batch Sampler, 117V Batch Sampler, 234V Gilson Interface Kit	523417 523418 514344
Auto 6 Cell Holder Options Unheated Water-regulated Peltier	523430 523415 523409
Peltier Temperature Controller 115V 230V	523422 523423
Cell Holders for 50 µL Microcells Micro-Auto 1 Micro-Auto 6	523382 523386
Cell Holder for 100 µL Multi-Microcell Micro-Auto 12	514254
Cell Holder for 5 µL Ultra-Microcell Adjustable Cell Holder	594439
Rectangular Cell Holder, 1 to 5 cm	517037
Cylindrical Cell Holder, 1 to 10 cm	517038

Gel Scan Accessories		
Gel Slits	596077	
Tube Holder	523428	
Film Holder	523429	
Supplies		
Dot Matrix Printer Supplies		
Fan Fold Paper	897516	
Ribbon Cartridge	596281	
X-Y Plotter Supplies		
Paper, 250 sheets	523274	
Pens, set of 4 (red, green, blue, black)	523298	
Cable (Male to female, RS-232	523198	
straight through, 3 meters.)		
Sipper Supplies		
Flowcell, 80 μL	599925	
Tubing Kit, Flowcell	514745	
Micro Sipper Cell, 50 μL	514758	
Tubing Kit, Micro Sipper Cell	514282	
Pump Tubing, Gray	651731	
Pump Tubing, Red	651767	
Waste Bottle	586656	
Waste Tubing	598549	
Microcells		
50 μL Microcell, 1 each	523270	
50 μL Microcell, 2 each	523450	
50 μL Microcell, 4 each	523451	
50 μL Microcell, 6 each	523452	
100 μL Multi-Microcell, 1 each, 4 position	523255	
Ultra-Microcells		
5 μL Ultra-Microcell with 100 capillaries	514261	
Replacement Capillaries, 100 each	514262	
Trace-Klean TM Solution		
Concentrated, 946 mL	589784	
Dilute, 473 mL	598190	

Replacement Parts

Fuses	
3A, slow blow, UL/CSA (for 110/120V instruments)	883908
1.6A, slow blow, IEC (for 220/240V instruments)	897176
UV Source	514366
Visible Source	945672

BECKMAN SALES AND SERVICE OFFICES

Beckman has local sales and service offices conveniently located throughout the world. For service, sales, or applications questions, contact the local Beckman office.

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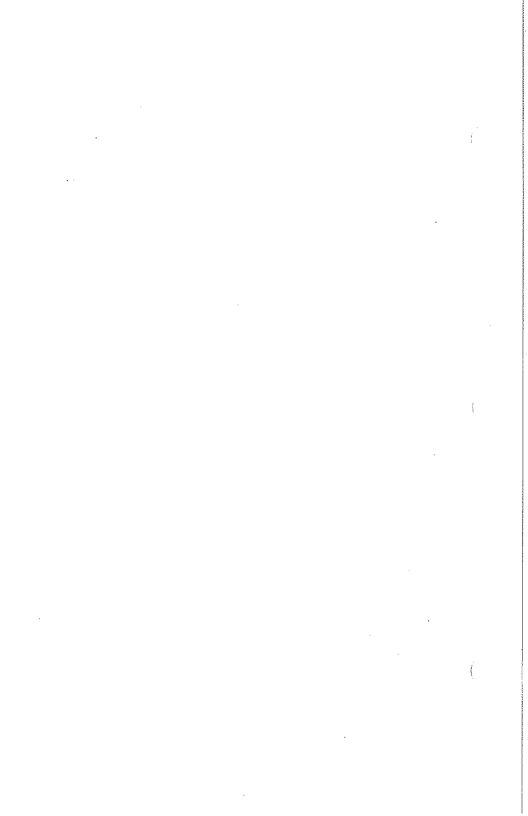
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Sales & Service 15-5



SECTION SIXTEEN

WARRANTY

Subject to the exceptions and upon the conditions stated below, Beckman warrants that this product shall be free from defects in workmanship and materials for one year after delivery of the products to the original Buyer by Beckman, and if any such product should prove to be defective within such one year period, Beckman agrees, at its option, either (i) to correct by repair or, at Beckman's election, by replacement with equivalent product any such defective product, provided that investigation and factory inspection discloses that such defect developed under normal and proper use, or (ii) to refund the purchase price. The exceptions and conditions mentioned above are as follows:

(a) components or accessories manufactured by Beckman which by their nature are not intended to and will not function for one year are warranted only to give reasonable service for a reasonable time; what constitutes reasonable time and reasonable service shall be determined solely by Beckman. A complete list of such components and accessories is maintained at the factory.

The following item is a consumable and has the following limited warranty:

514366 UV Source - 90 days from receipt

This limited warranty covers parts, only. Replacement instructions for this part are provided in section 12.2 of this manual. This limited warranty includes only the above named part.

- (b) Beckman makes no warranty with respect to components or accessories not manufactured by it; in the event of defect in any such component or accessory Beckman will give reasonable assistance to Buyer in obtaining from the respective manufacturer whatever adjustment is authorized by the manufacturer's own warranty.
- (c) any product claimed to be defective must, if required by Beckman, be returned to the factory, transportation charges prepaid, and will be returned to Buyer with transportation charges collect unless the product is found to be defective, in which case Beckman will pay all transportation charges.

1 6 (d) Beckman shall be released from all obligations under all warranties, either expressed or implied, if any product covered hereby is repaired or modified by persons other than its own authorized service personnel unless such repair by others is made with the written consent of Beckman.

IT IS EXPRESSLY AGREED THAT THE ABOVE WARRANTY SHALL BE IN LIEU OF ALL WARRANTIES OF FITNESS AND OF THE WARRANTY OF MERCHANTABILITY AND THAT BECKMAN SHALL HAVE NO LIABILITY FOR SPECIAL OR CONSEQUENTIAL DAMAGES OF ANY KIND OR FROM ANY CAUSE WHATSOEVER ARISING OUT OF THE MANUFACTURE, USE, SALE, HANDLING, REPAIR, MAINTENANCE OR REPLACEMENT OF THIS PRODUCT.

Representations and warranties made by any person, including dealers and representatives of Beckman, which are inconsistent or in conflict with the terms of this warranty, shall not be binding upon Beckman unless reduced to writing and approved by an expressly authorized officer of Beckman.

SECTION SEVENTEEN

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