2998 Photodiode Array Detector Operator's Guide

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Safety considerations

Some reagents and samples used with Waters instruments and devices can pose chemical, biological, and radiological hazards. You must know the potentially hazardous effects of all substances you work with. Always follow Good Laboratory Practice, and consult your organization's safety representative for guidance.

Safety advisories

Consult Appendix A for a comprehensive list of warning and caution advisories.

Operating this instrument

When operating this instrument, follow standard quality-control (QC) procedures and the guidelines presented in this section.

Applicable symbols

Symbol	Definition
Waters Corporation 34 Maple Street Milford, MA 01757 U.S.A.	Manufacturer
EC REP Waters Corporation Floats Road Withenshawe Manchester M23 9LZ United Kingdom	Authorized representative of the European Community
CE	Confirms that a manufactured product complies with all applicable European Community directives
ABN 49 065 444 751	Australia C-Tick EMC Compliant
c C used Us	Confirms that a manufactured product complies with all applicable United States and Canadian safety requirements
i	Consult instructions for use

Audience and purpose

This guide is intended for personnel who install, operate, and maintain the Waters 2998 Photodiode Array (PDA) detector.

Intended use of the 2998 Photodiode Array detector

Waters designed the 2998 Photodiode Array detector to analyze and monitor various types of compounds. The 2998 PDA detector is for research use only.

Calibrating

To calibrate LC systems, follow acceptable calibration methods using at least five standards to generate a standard curve. The concentration range for standards should include the entire range of QC samples, typical specimens, and atypical specimens.

When calibrating mass spectrometers, consult the calibration section of the operator's guide for the instrument you are calibrating. In cases where an overview and maintenance guide, not operator's guide, accompanies the instrument, consult the instrument's online Help system for calibration instructions.

Quality-control

Routinely run three QC samples that represent subnormal, normal, and above-normal levels of a compound. Ensure that QC sample results fall within an acceptable range, and evaluate precision from day to day and run to run. Data collected when QC samples are out of range might not be valid. Do not report these data until you are certain that the instrument performs satisfactorily.

ISM classification

ISM Classification: ISM Group 1 Class B

This classification has been assigned in accordance with CISPR 11 Industrial Scientific and Medical (ISM) instruments requirements. Group 1 products apply to intentionally generated and/or used conductively coupled radio-frequency energy that is necessary for the internal functioning of the equipment. Class B products are suitable for use in both commercial and residential locations and can be directly connected to a low voltage, power-supply network.

EC authorized representative



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1 2998 PDA Detector Optics Principles

To use the 2998 PDA detector effectively, you should understand the principles that underlie operation of the detector's optics and electronics.

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Detector optics

The detector is an ultraviolet/visible light (UV/Vis) spectrophotometer. With a photodiode array of 512 photodiodes and an optical resolution of 1.2 nm, the detector operates within a range of 190 to 800 nm.

The figure below illustrates the light path through the optics assembly of the detector.

Optics assembly light path



The following table describes the optics assembly components.

Optics assembly components

Component	Function
Lamp	Deuterium source lamp.
M1 mirror	Focuses light from the deuterium source lamp.

Component	Function	
Window	Used to help minimize air infiltration into the lamp housing.	
Filter flag/shutter	Flag positions for measuring open (sample) and blocked (dark) beam energies and a third for wavelength verification.	
Flow cell	Houses the segment of the flow path (containing eluent and sample) through which the polychromatic light beam passes.	
Spectrograph mirror and mask	The mirror focuses light transmitted through the flow cell onto the slit at the entrance to the spectrographic portion of the optics. The mirror mask defines the size of the beam at the grating.	
Slit	Determines wavelength resolution and intensity of light striking the photodiodes. The width of the slit is 50 µm.	
Grating	Disperses light into bands of wavelengths and focuses them onto the plane of the photodiode array.	
Order filter	Reduces the contribution of second-order diffraction of UV light (less than 370 nm) to the light intensity observed at visible wavelengths (greater than 370).	
Photodiode array	A linear array of 512 photodiodes. The diode width (50 μm), together with a 50-μm slit, yields single wavelength resolution of 1.2 nm.	

Optics assembly components (Continued)

Calculating absorbance

The detector computes absorbance by subtracting the dark current (see "Dark current" on page 1-9) from the reference spectrum (reference energy) and the acquired spectrum (sample energy). Absorbance is based on the principles of Beer's law.

Beer's law

The Beer-Lambert law (commonly called Beer's law) describes the relationship between the quantity of light of a particular wavelength arriving

at the photodiode and the concentration of the sample passing through the flow cell. Beer's law is expressed as $A = \varepsilon lc$ where

A = dimensionless quantity measured in absorbance units

- ϵ = constant of proportionality known as the molar absorptivity
- l = path length in centimeters (1.0 cm in the detector's normal flow cell)
- c =concentration in moles per liter

Beer's law applies only to well-equilibrated dilute solutions. It assumes that the refractive index of the sample remains constant, that the light is monochromatic, and that no stray light reaches the detector element. As concentration increases, the chemical and instrumental requirements of Beer's law are sometimes violated, resulting in a deviation from (absorbance versus concentration) linearity. The absorbance of mobile phase can reduce the linear range.

Absorbance as a function of concentration



Flow cell operating principles

The Waters TaperSlit[™] flow cell used in the 2998 detector renders the detector baseline essentially insensitive to changes in mobile phase refractive index (RI). RI changes occur during gradient separations or result from temperature or pump-induced pressure fluctuations.

To achieve RI immunity, a combination of a spherical mirror, a lens at the entrance of the flow cell, and a taper to the internal bore of the flow cell

prevents light rays from striking the internal walls of the flow cell. The Waters TaperSlit flow cell, so-called because of the shape of the flow cell exit face, matches the shape of the spectrograph slit. Compared to a conventional flow cell with a cylindrical shape, the 2998 detector achieves higher light throughput for a given spectral resolution with the TaperSlit cell design.



Comparison of flow cell characteristics

Resolving spectral data

Together with photodiode spacing, the detector's $50-\mu m$ slit determines the intensity and bandwidth of the light that strikes the photodiode array. Reducing the bandwidth increases the resolving power of the detector. As a result, similar spectra are more effectively distinguished.

The grating images the slit onto the photodiode array. The angle of diffraction from the grating determines the wavelength that strikes a particular photodiode in the array.

The following figure shows an absorbance spectrum of benzene. Note that the wavelength resolution is sufficient to resolve five principal absorption peaks.

Benzene spectrum at different resolutions



Measuring light at the photodiode array

The detector measures the amount of light striking the photodiode array to determine the absorbance of the sample in the flow cell.

The array consists of a row of 512 photodiodes. Each photodiode acts as a capacitor initially holding a fixed amount of charge.

Light striking a photodiode discharges the diode. The magnitude of the discharge depends on the amount of light striking the photodiode.

Photodiodes discharged by light



The detector measures the amount of charge required to recharge each photodiode. The charge is proportional to the amount of light transmitted through the flow cell over the interval specified by the diode exposure time.

Optimizing the signal-to-noise ratio

To optimize signal-to-noise ratios, choose an acquisition wavelength range that includes only the wavelengths of interest. Ensure that the range is one in which the mobile phase absorbs only minimally (see Appendix C). You can further improve the signal-to-noise ratio by increasing the bandwidth parameter. For example, you can choose to operate at 3.6 nm instead of at 1.2 nm resolution. See page 1-10 for more information.

Optimizing filter constants

The filtering constant you select affects the peak intensity. To increase sensitivity, decrease the filter time constant.

Selecting the appropriate sampling rate

A sufficient number of points must fall across a peak to define its shape. Thus, the definition between peaks is lost at very low sampling rates. Empower uses

the end time minus the start time, to calculate the Points Across Peak value for each integrated peak in the chromatogram.

Tip: The Points Across Peak value appears in the Peaks table, at the bottom of the Review Main window. If the Points Across Peak field is not visible, right-click anywhere in the table, and then click Table Properties. Click the Columns tab, and then scroll down to find the Points Across Peak field. Clear the check box, and then click OK.

If the Points Across Peak value for the narrowest peak of interest is less than 25, you should specify a higher sampling rate in the instrument method. If the value is greater than 50, you should specify a lower sampling rate in the instrument method.

Example of how baseline noise increases with higher sampling rates



Computing absorbance data points

The absorbance data points are calculated by the detector and then transmitted to the database (MassLynx or Empower).

The absorbance calculation is

$$Absorbance_{t,\lambda} = Log_{10} \left[\frac{R_{\lambda} - D_{\lambda}}{S_{t,\lambda} - D_{\lambda}} \right]$$

where

S = sample energy

D = dark energy

R = reference energy

t = elapsed time after start of injection

 $\lambda = wavelength$

This value is then filtered as specified.

Dark current

Photodiodes discharge even when they are not exposed to light. This discharge is called dark current. The dark current is updated by closing the shutter to take a dark current reading for each diode. After updating, the detector opens the shutter and then subtracts the dark current values as shown in the above equation.

Reference spectrum

The reference spectrum is a measure of lamp intensity and mobile phase absorbance at initial conditions. The detector records a reference spectrum at the start of each injection. The reference spectrum value is calculated using the same filter time constant as the absorbance data.

Data averaging

The data the detector reports to the database (Empower or MassLynx) can be the average of a number of data points. After calculating absorbance, the detector averages absorbance values based on the requested spectral resolution.

Averaging spectral resolution

The detector can simultaneously collect two types of data channels: spectra (3D) and chromatograms (2D). For best results in library matching and peak purity analysis, set the 3D resolution to 1.2 nm.

For chromatograms (2D data), select a resolution to optimize signal amplitude, baseline noise, and linear dynamic range. When the monitoring wavelength for an analyte corresponds to a peak's lambda max, increasing bandwidth tends to decrease peak height while reducing baseline noise and linear dynamic range.

Tip: A resolution of 3.6 nm is effective for many analytes.



Resolution comparison for anthracene

Averaging chromatographic sampling rate

Sample rate is the number of data points acquired per second. The number of times a given pixel is read during the sample rate interval depends on its exposure time. For example, if exposure time is 25 msec and sample rate is 20 Hz, then exposures per sample is

20 samples/second = (1000 msec/second) (20 samples/second)(25 msec/exposure) = 2 exposures/sample The readings are averaged and reported as a single data point. The baseline noise decreases as the number of exposures/sample increases.

Tip: The amount of data stored is based on wavelength range, spectral resolution, runtime, and sample rate. Specify these parameter values in the General tab of the PDA Instrument Method Editor. For details, refer to the Empower or MassLynx online Help.

Filtering data

In the General tab of the PDA Instrument Method Editor (for details, refer to the Empower or MassLynx online Help) you can apply an optional noise filter (via the Digital Filtering parameter) to the data acquired. The following table lists the digital filter settings for the allowable data rates.

Data Rate (Hz)	Slow (sec)	Normal (sec)	Fast (sec)	Range (sec)	Increment (sec)
1	4.000	2.000	1.000	No filter, 0.100 to 5.000	0.100
2	2.000	1.000	0.500	No filter, 0.100 to 5.000	0.100
5	0.800	0.400	0.200	No filter, 0.100 to 2.000	0.100
10	0.400	0.200	0.100	No filter, 0.100 to 1.000	0.100
20	0.200	0.100	0.050	No filter, 0.050 to 0.500	0.050
40	0.100	0.050	0.025	No filter, 0.025 to 0.250	0.025
80	0.050	0.025	0.0125	No filter, 0.0125 to 0.100	0.0125

Digital filter settings for data rates

Tip: For best chromatography resolution, choose Fast to avoid peak broadening.

Reference wavelength compensation

A compensated reference wavelength collects wide-band absorbance data in a region of the spectra where no known analytes exist. It is used to reduce detector drift and wander, which can affect integration quality.

The detector calculates the compensation value by averaging the absorbance values within a range of wavelengths that you select. It then subtracts that value from the absorbance value:

Abs-Comp(t) = Abs(t) - CRef(t)

where

Abs-Comp = absorbance-compensated

Abs = absorbance

CRef = compensation reference

t = elapsed time after start of injection

A starting and ending wavelength defines the compensation reference. CRef bandwidth must be ≥ 40 nm and ≤ 100 nm, and fall within a range of 190 to 800 nm.

Tip: Select a compensation reference range where no analytes are expected to appear. Since responses are subtracted from the absorbance value, any responses within the compensation reference range can erroneously affect quantitative data.

2 Setting Up the Detector

The chapter provides the information necessary to set up the detector.

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Contents:

Before you begin

Requirement: To install the 2998 PDA detector, you should know how, in general, to set up and operate laboratory instruments and computer-controlled devices and also how to handle solvents.

Before installing the detector, ensure that

- the required components are present.
- none of the shipping containers or unpacked items are damaged.

Unpacking and inspecting

The detector shipping carton contains:

- Certificate of Structural Integrity
- 2998 PDA detector
- Waters 2998 PDA Detector Operator's Guide (this document)
- Startup kit
- Release notes

To unpack the detector and flow cell:

- 1. Check the contents of the shipping cartons against the packing lists to ensure that you received all items.
- 2. Save the shipping cartons for future transport or shipment.

If you discover any damage or discrepancy when you inspect the contents of the cartons, immediately contact the shipping agent and your local Waters representative.

Customers in the USA and Canada should report damage and discrepancies to Waters Technical Service (800 252-4752). Others should phone their local Waters subsidiary or Waters corporate headquarters in Milford, Massachusetts (USA), or visit http://www.waters.com and click Offices.

For complete information on reporting shipping damages and submitting claims, see *Waters Licenses, Warranties, and Support Services*.

Make sure the instrument serial number on the rear panel nameplate or inside the front panel corresponds to the number on the instrument validation certificate.

Laboratory site selection

To ensure the reliable operation of your detector:

- do not situate it under a heating or cooling vent
- connect it to a power supply that is grounded AC, 100 to 240 VAC
- provide clearance of at least 15 cm (6 inches) on the back side for ventilation



Caution: To avoid damaging the detector, the weight of items stacked on top of it should not exceed 18.1 kg (40 pounds).

The dimensions shown in the figure "Detector dimensions" on page 2-4 indicate that the detector is a stackable unit and does not require extra bench space.

Place the detector close to the outlet of the column to minimize band broadening, which will reduce the resolution of the chromatogram.

Requirement: Mount the detector on a level surface to allow proper function of the drip management system (drain tube), to which you can connect a waste reservoir that diverts solvent leaks from the flow cell.

Detector dimensions

The following figure shows the dimensions of the detector.

Detector dimensions



Connecting to the electricity source

The 2998 PDA detector requires a separate, grounded electricity source. The ground connection in the electrical outlet must be common and connected near the system.



Warning: Avoid electrical shock:

- Use power cord SVT-type in the United States and HAR-type or better in Europe. For other countries, contact your local Waters distributor.
- Power-off and unplug the detector before performing any maintenance on the instrument.
- Connect the detector to a common ground.

To connect to the electricity source:

Recommendation: Use a line conditioner or an uninterruptible power supply (UPS) for optimum long-term input voltage stability.

- 1. Place the Off/On (O/I) switch in the Off (O) position.
- 2. Connect the female end of the power cord to the receptacle on the rear panel of the detector.
- Connect the male end of the power cord to a suitable wall outlet.
 Important: Do not turn the detector on at this time.

Making signal connections

See also: Waters Ethernet Instrument Getting Started Guide.

The following figure shows the rear panel location of the connectors used to operate the detector with external devices.

2998 PDA Detector rear panel



The signal connections you need to make to your detector depend on the signal connections available on the other instruments in your HPLC system.

Connecting the Ethernet cable

A Waters instrument communicates with the acquisition computer through the dedicated local area network (LAN). At the acquisition computer, the instrument network card provides the interface that makes communication possible.

You must install the Waters instrument software driver in the acquisition computer so that the computer can control the instrument. See the software installation instructions that accompany the instrument control software.

Single Waters instrument connection

In a single Waters instrument system configuration, the connection hardware requires only one standard Ethernet cable provided in the startup kit.

Single Waters instrument connection



Multiple Waters instrument connections

System configurations with many Waters Ethernet instruments require an Ethernet switch to communicate multiple signals among Waters instruments and the acquisition computer.

Connection hardware requires one standard Ethernet cable per Waters instrument and a standard Ethernet cable to connect the network switch and the acquisition computer.

You must install the Waters instrument control software in the acquisition computer so that the computer can control the Waters instrument. See the software installation instructions that accompany the software instrument driver disk.

Network installation guidelines

Configurations for multiple Waters instruments use a dedicated LAN, which requires a design based on the following guidelines:

- Ethernet cable
- A maximum distance of 100 meters (328 feet)

Requirement: You must use a network switch for multiple Ethernet instruments. Network hubs are not supported.

Waters Ethernet instrument connections



Making inject start signal connections

When you are using an Ethernet data system with the detector, the data system or controller must receive an inject-start signal from the autosampler or manual injector to initiate the data collection and time-based programs. The following table summarizes the inject start connections for different system configurations.

Detector inject start connections

Inject start output source	Inject start input connection (on the 2998 PDA detector, connector A)
Waters Alliance Separations Module	Inject Start + / –
Waters 717 Autosampler	Inject Start + / –
Waters manual injector, or third-party manual injector or autosampler	Inject Start + / –

Tip: If the injector is an e2695 separations module running in Ethernet mode or a 2707 autosampler, then the inject start cable should not be connected. However, if the injector is an e2695 separations module running in IEEE mode, then the inject start cable should be connected.

Connecting to a manual injector

If you are using a manual injector with your system, connect the signal cables from the rear panel connector on the detector to the injector as shown in the following table.

Detector connections to a manual injector

2998 PDA detector (connector A)	Manual injector
Inject Start + (red)	One set of spade lug Inject Start
Inject Start – (black)	terminals

For information on injection trigger signals from a manual injector, see page 2-15.

Connecting to other instruments

This section describes signal connections between the detector's rear panel and these items:

- Waters Alliance Separations Module
- Waters 1500-series pump

- Waters SAT/IN[™] Module
- Waters, or other, manual injector
- Waters autopurification system
- Other manufacturers' integrator or A/D interface device

Warning: To avoid electrical shock, power-off instruments being connected before making any electrical connections.

Requirement: To meet the regulatory requirements of immunity from external electrical disturbances that can affect the performance of this instrument, do not use cables longer than 3 meters (9.8 feet) when you make connections to the I/O connectors. In addition, ensure you connect the shield of the cable to ground at one instrument only.

To connect the detector to other instruments, use the two analog-out/event-in (I/O) connectors and their mating connectors on the rear panel.

2998 rear panel analog-out/event-in connectors



The following table describes the detector I/O connections.

Signal connections	Description
Inject start	Activates timed events by triggering the run-time clock to start.
Lamp off	When the input is enabled, the lamps is turned off. The lamp can only be turned on by sending a new method to the detector, using the lamp button, or rebooting the detector.
Chart mark	Adds a chart mark (at 10% of full scale) to either or both analog output channels (Signal Out 1 and Signal Out 2) and is configurable.
Auto zero	Calculates an offset value that, when added to the sample signal, makes the resulting baseline signal zero.
Analog 1 and Analog 2	Method programmable analog output. Minimum output voltage range: -0.1 to 2.1 VDC For sample rates of 10, 20, 40, or 80 Hz, this output runs at the selected data rate. For sample rates of 1, 2, or 5 Hz, this output runs at 10 Hz.
Switch 1	Controls a timed event or threshold level and is a user-programmable auxiliary output.
Switch 2	Controls a timed event or threshold level and is a user-programmable auxiliary output.

Detector analog-out/event-in connections

Generating inject start

To generate the inject start function on the detector at the start of an injection from the Alliance Separations Module, make the connections shown in the following table and figure.
Tip: The firmware defaults to auto zero on inject.

Detector connections to an Alliance separations module

Alliance separations module (connector B)	2998 PDA detector (connector A)
Pin 1 inject start (red)	Pin 1 inject start + (red)
Pin 2 inject start (black)	Pin 2 inject start – (black)

Inject start connections between the Alliance separations module and the detector



Generating stop flow

Warning: Fire and explosion hazard. Failure to connect the stop flow output could cause the detector to flood.

The detector has a programmable switch output, that can be controlled by threshold or timed events.

To generate stop flow, make the connections shown in the following table and figure.

Requirement: To automatically stop the chromatographic flow to the system in the event of an error condition or hardware failure, the stop flow signal must be connected to the chromatographic pump.

Detector connections to an Alliance separations module

Alliance separations module (connector B)	2998 PDA detector (connector B)
Pin 4 stop flow + (red)	Pin 6 switch 1 + (red)
Pin 5 stop flow – (black)	Pin 7 switch 1 – (black)

Stop flow connections between the Alliance separations module and the detector



2998 PDA detector connector B

Connecting to a chart recorder or external analog data collection device

To send an analog output signal from the detector to a chart recorder, make the connections shown in the following table and figure.

Analog output connections to a chart recorder

Chart recorder connectors	2998 PDA detector (connector B)
Pen 1 +	Pin 1 signal out + (red)
Pen 1 – Pin 2 signal out – (black)	
Tip: For this connection, do not use the cable shield.	

Analog output connections to a chart recorder



Connecting to an Empower or MassLynx data system using an eSAT/IN module

To send an integrator analog output signal (-0.1 to +2.1 V) from the detector to an Empower or MassLynx system (through a two-channel SAT/IN Module), make the connections shown in the following table and figure.

Detector connections to the eSAT/IN module

SAT/IN module connector	2998 PDA detector (connector B)
CHANNEL 1	Pin 1 signal out + (white)
	Pin 2 signal out – (black)

Analog output connections to the eSAT/IN module



Connecting injection trigger signals

The detector accepts the following injection trigger signals from a manual injector:

- Inject start signal from a contact closure signal with each injection
- Auto zero signal to adjust the zero offset of the detector each time the injector makes an injection

Each time the detector receives a signal from an injector, it performs the corresponding auto zero or inject start function.

To send an auto zero or chart mark signal from an injector to the detector, make the connections shown in the following tables and figures.

Tip: The firmware defaults to auto zero on inject.

Inject start connections to an injector (pulse duration 0 to 10 seconds)

2998 PDA detector (connector A)	Injector connector
Pin 1, inject start + (red)	Two spade lug (or similar) terminal
Pin 2, inject start – (black)	connectors.

Inject start connections to an injector



Plumbing the detector

See also: "Minimum tubing bend radius recommendations" on page C-6.

Warning:

- To avoid chemical hazards, always observe Good Laboratory Practices when operating your system. Refer to the Material Safety Data Sheets shipped with solvents for handling information.
- Using incompatible solvents can cause severe damage to the instrument and injury to the operator.



Caution: To prevent contamination, wear powder-free, nonlatex gloves when plumbing the detector.

Required Materials

- 5/16-inch open-end wrench
- 0.009-inch (0.23 mm) ID stainless steel tubing (included in the Startup Kit)
- Stainless steel tubing cutter or scribing file
- Pliers, plastic-, or cloth-covered
- Compression screw assemblies (3)

To make plumbing connections to the detector:



Caution: To avoid damaging the flow cell, never cut the 1/32-inch stainless steel flow cell tubing.

1. Measure the length of tubing needed to connect the column outlet to the detector inlet.

Tip: Keep the length of this tubing as short as possible to prevent band broadening.

2. Measure the length of tubing needed to connect the detector outlet to a waste collection bottle.

Tip: Ensure the length of this tubing is at least 30 to 60 cm (1 to 2 feet) to prevent air bubbles from forming in the flow cell.

3. Cut the two lengths of tubing as follows:

- a. Use a Waters 1/16-inch stainless steel tubing cutter or a file with a cutting edge to scribe the circumference of the tubing at the desired break point.
- b. Grasp the tubing on both sides of the scribed mark with cloth- or plastic-covered pliers (to prevent marring the surface), then gently work the tubing back and forth until it separates.
- c. File the tubing ends smooth and straight to minimize dead volume and band broadening.
- 4. Assemble a reusable finger-tight fitting at both ends of the column outlet line and at one end of the detector outlet line.



Reusable finger-tight fitting assembly

- 5. Seat one end of the column outlet tubing in the fitting of the column outlet, finger-tighten the reusable finger-tight fitting until it is snug, and then add a quarter-turn.
- 6. Seat the other end of the tubing in the fitting of the detector inlet, then tighten the reusable finger-tight fitting as in step 5.
- 7. Seat the end of the detector outlet tubing with the reusable finger-tight fitting in the detector outlet fitting, finger-tighten the fitting until it is snug, and then add a quarter-turn. Insert the other end of the tubing in the waste container.

0

Caution: To avoid damage to the flow cell, avoid pressures approaching its maximum allowable pressure, 6895 kPa (69 bar, 1000 psi) (70 kg/cm²).

8. Slide the 3/8-inch OD Tygon[®] tubing, supplied in the startup kit, over the barbed drain fitting located on the drip tray and route it to a suitable waste container.

Making the gas supply connection

The detector can be connected to a nitrogen source to improve operation at lower wavelengths.

Requirement: Only use a constant supply 28 to 41 kPa (.28 to .41 bar, 4 to 6 psi) of dry, oil-free, filtered nitrogen.

Warning: Do not use gases that would allow the combustion of combustible solvents. Always use inert gases.

To make the gas supply connection:

1. Connect one end of the plastic tubing to the nitrogen source.

Tip: The nitrogen source may require an adapter in order to fit the plastic tubing.

2. Connect the other end of the nitrogen fitting on the rear panel of the detector.

Starting up and shutting down the detector



Warning:

- Always observe Good Laboratory Practices when you use this equipment and when you work with solvents and test solutions. Know the chemical and physical properties of the solvents and test solutions you use. See the Material Safety Data Sheet for each solvent and test solution in use.
- Using incompatible solvents can cause severe damage to the instrument and injury to the operator.

The detector startup procedure takes less than one minute. Once completed, you should allow the detector to warm up for at least one hour before running an analysis. Follow the procedures in this section to ensure reliable detector performance.

Starting up the detector

To start up the detector:

1. In your instrument method, set the solvent delivery system or pump to deliver 10 mL of HPLC-grade water. For details, refer to the Empower or MassLynx online Help.

Guidelines:

- Use only thoroughly degassed HPLC-grade solvents. Gas in the mobile phase may form bubbles in the flow cell and cause the detector to fail the reference energy diagnostic test.
- Ensure that the priming solvent is correctly composed and that it is of high quality and miscible with any other solvents used in your system. Use filters in all solvent reservoirs, and ensure the volumes of solvents are sufficient for priming.
- 2. Press the Off/On (O/I) switch on the front panel of the detector to the On (I) position.
- 3. Observe the Lamp and Power indicator LEDs on the front panel of the detector.

The power and lamp LEDs change as follows:

- The power LED shows green.
- During initialization, the power LED flashes green.
- After the detector has successfully powered-on, the power and lamp LEDs show steady green.

4. Wait 1 hour for the detector to stabilize before acquiring data. If the detector fails to stabilize, see Chapter 4.



Detector indicator lights

Monitoring detector LEDs

Light-emitting diodes on the detector indicate its state of functioning.

Power LED

The power LED, on the detector's front panel, indicates when the instrument is powered-on or powered-off. It is steady green when the instrument is working properly.

Lamp LED

The lamp LED, to the left of the power LED, indicates the lamp status.

Warning: To avoid electric shock, do not touch the lamp connector if the lamp LED is flashing red. Power-off and unplug the detector before touching the lamp connector.

The following table identifies each LED mode with its corresponding detector lamp state.

LED mode and color	Description
Unlit	Indicates the detector lamp is extinguished.
Constant green	Indicates the detector lamp is ignited.
Flashing green	Indicates the detector is initializing or verifying calibration.
Flashing red	Indicates that an error stopped the detector. Refer to the console for information regarding the error.
Constant red	Indicates a detector failure that prevents further operation. Power-off the detector, and then power-on. If the LED is still steady red, contact your Waters service representative.

Lamp LED indications

Shutting down the detector

To shut down the detector:

- 1. If the mobile phase contains buffers, set the solvent delivery system or pump to deliver 10 mL of HPLC-grade water. Otherwise, set the solvent delivery system or pump to deliver 10 mL of degassed methanol.
- Press the Off/On (O/I) switch on the front panel of the detector to the Off (O) position.

Using a cuvette

The detector cuvette option allows for ease of use in:

- Sample handling
- Instrument verification and qualification

The detector uses a standard 10-mm path length spectrophotometric cell (quartz cuvette). You insert the cuvette with a frosted side facing up in the cuvette holder, which you then place in the detector flow cell assembly.

2998 PDA Detector cuvette holder with the cuvette inserted



Restriction: Because the measurement is actually a composite of both the contents of the cuvette and the flow cell, you need to perform cuvette measurements under identical flow cell conditions. If you store spectra and acquire new spectra for subtraction, you need to be aware of differences, if any, in flow cell conditions.

Ideally, you should perform both the zero and sample measurements using the cuvette when the HPLC instruments are in the idle or static state and operating under identical flow cell conditions.

Caution: Handle the cuvette gently and on the frosted sides only. Fingerprints on the clear quartz interfere with the light path and compromise the integrity of cuvette measuring operations.

Before you begin

Recommendation: To ensure accurate results, use a 10-mm path length quartz cuvette and matched pairs (from the same manufacturing lot) of quartz cuvettes for your zero and sample measurements.

Before beginning a measurement using the cuvette:

- 1. Fill the flow cell by flushing with 10 mL of the same mobile phase that you are going to use with the cuvette measurement.
- 2. To ensure a clear and clean cuvette, wipe the clear portion of the cuvette with low-lint, nonabrasive tissue wipes.

Cuvette measuring procedure

To begin a cuvette measurement:

- 1. Remove the detector front left panel cover.
- 2. To remove the cuvette holder, slide it toward you.



3. With the spring guide facing you, gently insert the cuvette (containing eluent) up and under the guide, with the cap facing upward (into the holder) and a frosted side of the cuvette facing up. Refer to the figure on page 2-22.

Recommendations:

- Ensure that you have enough liquid (3 mL) in the cuvette so that, when it is inserted into the holder, you can see liquid through the cuvette holder aperture (that is, the liquid completely covers the aperture).
- Because the cuvette holder is angled, use your thumb or forefinger to ensure the cuvette is secure in the slot and does not slide forward.
- Ensure that it does not become dislodged when you are replacing the cuvette holder.
- 4. Gently guide the cuvette holder back into the flow cell assembly until it sits in position securely.

- 5. Reinstall the detector front left panel cover:
 - To prevent invalid subsequent chromatographic results, remove the cuvette from the detector, and replace the empty holder after running your cuvette measurements.
 - To maintain optimum system performance, replace the front left panel cover before resuming normal operation of the detector.
- 6. Insert a reference cuvette containing the mobile phase standard, and run a zero measurement.
- 7. Replace the reference cuvette with a cuvette containing your analyte dissolved in mobile phase solvent, and run a sample measurement.
- 8. Use the storage, review, subtract and review, and replay functions to analyze the data obtained.

3 Maintaining the Detector

This chapter describes the routine maintenance procedures you should perform on the detector.

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Maintenance considerations	3-2
Routine Maintenance	3-3
Maintaining the flow cell	3-4
Replacing the lamp	3-7
Replacing the fuses	3-10

Contacting Waters technical service

If you are located in the USA or Canada, report malfunctions or other problems to Waters Technical Service (800 252-4752). Otherwise, phone the Waters corporate headquarters in Milford, Massachusetts (USA) or contact your local Waters subsidiary. Our Web site includes phone numbers and e-mail addresses for Waters locations worldwide. Go to www.waters.com, and click About Waters > Worldwide Offices.

When you contact Waters, be prepared to provide this information:

- Nature of the symptom
- Instrument serial number
- Solvent(s)
- Method parameters (sensitivity and wavelength)
- Type and serial number of column(s)
- Sample type
- Empower or MassLynx software version and serial number

For complete information on reporting shipping damages and submitting claims, see *Waters Licenses, Warranties, and Support Services*.

Maintenance considerations

Safety and handling

Observe these warning and caution advisories when you perform maintenance on your detector.

Warning: To prevent injury, always observe Good Laboratory Practices when you handle solvents, change tubing, or operate the system. Know the physical and chemical properties of the solvents you use. See the Material Safety Data Sheets for the solvents in use.



Warning: Avoid electric shock:

- Do not open the detector cover. The detector does not contain user-serviceable components.
- Power-off and unplug the detector before performing any maintenance on the instrument.



Caution: To avoid damaging electrical parts, never disconnect an electrical assembly while power is applied to the detector. To interrupt power to the detector, set the power switch to Off, and then unplug the power cord from the AC outlet. Wait 10 seconds before you disconnect an assembly.

Spare parts

Waters recommends that you replace only parts mentioned in this document. For spare parts details, see the Waters Quality Parts Locator on the Waters web site's Services/Support page or the *2998 PDA Detector spare parts list* (part number 71500121906).

Routine Maintenance

The 2998 PDA detector requires minimal routine maintenance.

To achieve optimal performance, follow these steps:

- 1. Replace solvent reservoir filters in your HPLC system regularly.
- 2. Filter and degas solvents to prolong column life, reduce pressure fluctuations, and decrease baseline noise.
- 3. Flush buffered mobile phases out of the detector with HPLC-grade water followed by a 5 to 10% methanol solution each time the detector is shut down. This process prevents:
 - Plugging of the solvent lines and flow cell
 - Damage to the components
 - Microbial growth

Maintaining the flow cell

The flow cell requires maintenance when

- the reference spectrum changes.
- the cell fluid leaks out of the drain tube.
- the detector cannot initialize but the lamp is in good condition.
- the detector causes high backpressure.

Tip: Conditions other than a dirty flow cell can cause decreased lamp intensity. For more information, refer to Chapter 4.

Flow cell maintenance consists of

- flushing the flow cell.
- removing the flow cell.
- disassembling and cleaning the flow cell.
- installing the flow cell assembly.

Flushing the flow cell

Required Materials

- HPLC-grade water
- HPLC-grade methanol

If the flow cell requires cleaning, first try flushing it with solvent.

To flush the flow cell:

1. Select a solvent compatible with the samples and mobile phases that you have been using. If you have been using buffers, flush with 10 mL of HPLC-grade water, then flush with 10 mL of a low-surface-tension solvent such as methanol.

Tip: Ensure that the solvent is miscible with the previous mobile phase.

2. Test the lamp energy by performing the Read energy diagnostic test (see page 4-3).

If the lamp diagnostic test fails and the lamp has not been used more than 2000 hours or 1 year from date of purchase (whichever comes first), call Waters Technical Service (see page 3-2).

Replacing the flow cell

Required materials

- 1/4-inch flat-blade screwdriver
- Flow cell

To replace the flow cell:

- 1. Power-off the detector.
- 2. Stop the solvent flow.
- 3. Lift up the detector front cover and pull the front cover from the detector chassis.
- 4. Disconnect the detector's inlet tubing from the column outlet connection.

Lamp Flow cell handle Thumbscrews

2998 PDA detector analytical flow cell

Tip: The optional auto-purification flow cell has three fittings – inlet analytical, inlet prep, and outlet.

5. Remove the flow cell:

- Use a 1/4-inch flat-blade screwdriver to loosen the three thumbscrews on the flow cell assembly's front plate.
- Grasp the handle and gently pull it toward you.
- 6. Unpack and inspect the new flow cell.
- 7. Square the flow cell assembly in front of the opening, and then insert it into the optics bench. Note that the flow cell uses the alignment pins on the optics bench.
- 8. Gently push the front of the assembly until it seats on the front alignment pins.

Alignment pin

Installing the analytical flow cell assembly

9. Continue to insert the flow cell until the three thumbscrews align with their holes in the bulkhead.

Caution: To prevent the flow cell from binding and ensure that it is properly seated in the bulkhead, alternate between tightening the captive screws and pushing the flow cell forward.

- 10. Hand tighten the thumbs crews, and then tighten them snug with the 1/4-inch flat-blade screwdriver.
- 11. Connect the inlet tubing to the column outlet connection and flow cell inlet.
- 12. Replace the front panel cover.
- 13. Before you power-on the detector, prime the system to fill the flow cell with solvent and remove any air.

Replacing the lamp

Change the lamp when it repeatedly fails to ignite or when the detector fails to calibrate.

The 2998 detector source lamp is warranted to light and pass startup diagnostic tests for 2000 hours or 1 year from the date of purchase, whichever comes first.



Warning: To prevent burn injuries, allow the lamp to cool for 30 minutes before removing it. The lamp housing gets extremely hot during operation.



Warning: To avoid eye injury from ultraviolet radiation exposure,

- power-off the detector before changing the lamp.
- wear eye protection that filters ultraviolet light.
- keep the lamp in the housing during operation.

To remove the lamp:

- 1. Power-off the lamp.
- 2. Power-off the detector and disconnect the power cable from the rear panel.
- 3. Allow the lamp to cool for 30 minutes.

Warning: The lamp and lamp housing can be hot. Wait 30 minutes after powering off the detector for these components to cool before touching them.

4. Lift up the front panel cover and pull it away from the chassis.

5. Detach the lamp power connector from the detector.



Removing the lamp

Warning: Lamp gas is under slight negative pressure. To prevent shattering the glass, use care when disposing of the lamp. Waters suggests that you adequately cushion an old lamp by containing it in the packaging of its replacement before you dispose of it.

6. Loosen the two captive screws in the lamp base. Gently withdraw the lamp from the lamp housing.

To install the lamp:

Caution: Do not touch the glass bulb of the new lamp. Dirt or fingerprints adversely affect detector operation. If the bulb needs cleaning, gently rub it with ethanol and lens tissue. Do not use abrasive tissue. Do not apply excessive pressure.

1. Unpack the new lamp from its packing material without touching the bulb.

- 2. Inspect the new lamp and lamp housing.
- 3. Position the lamp so that the cut-out on the lamp base plate is at the 1 o'clock position, in line with the alignment pin on the lamp housing, and then gently push the lamp forward until it bottoms into position. Ensure that it is flush to the optics bench.



Caution: To prevent the lamp from binding and ensure that it is properly seated in the lamp housing, alternate between tightening the captive screws and pushing the lamp forward.

- 4. Tighten the two captive screws, and then reconnect the lamp power connector.
- 5. Replace the front panel cover.
- 6. Power-on the detector, and then wait 1 hour for the lamp to warm before resuming operations.

Tip: Cycling power to the detector (that is, powering-off and then powering-on the instrument) initiates the verification procedures.

7. In the console, select Maintain > Change Lamp.

Change Lamp dialog box

Cł	iange Lamp				
	Currently Insta	led Lamp			
	State:	On an	d stable		
	Good ignitior	is: 28			
	Total ignition	is: 28			
	Date Installed	Serial Number		Hours	
		(Current)		80.20	
					_
					_
		New Lamp	Print	Close	

8. Click New Lamp.

New Lamp dialog box

New Lamp	
Serial number of new la	imp:
ОК	Cancel

9. Type the serial number for the new lamp (see the label attached to the lamp connector wire), and then click OK.

Replacing the fuses



The detector requires two 100 to 240 VAC, 50 to 60-Hz, F 3.15-A, 250-V (fast-blow), 5×20 mm (IEC) fuses.

Suspect a fuse is open or otherwise defective when

- the detector fails to power-on.
- the fan does not operate.

To replace the fuses:

Requirement: Replace both fuses, even when only one is open or otherwise defective.

- 1. Power-off the detector and disconnect the power cord from the power entry module.
- 2. Pinch the sides of the spring-loaded fuse holder, located above the power entry module on the rear panel of the detector.
- 3. With minimum pressure, withdraw the spring-loaded fuse holder.

Removing the fuseholder



- 4. Remove and discard the fuses.
- 5. Make sure that the new fuses are properly rated for your requirements, and then insert them into the holder and the holder into the power entry module, gently pushing until the assembly locks into position.
- 6. Reconnect the power cord to the power entry module.

4 Diagnostic Tests and Troubleshooting

Consult this chapter when troubleshooting problems with the 2998 PDA detector. However, bear in mind that the detector measures only the bulk properties of a system. Therefore, the cause of an apparent detector problem may actually originate with the chromatography or other system instruments.

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Diagnostic tests

The 2998 PDA detector automatically runs a series of internal diagnostic tests upon start up. The indicator LEDs on the front of the detector and messages at the Empower workstation show the results of the start up internal diagnostic tests.

If you need to determine the cause of a problem during operation of the detector, you can run the same internal diagnostic tests from the Empower workstation. Additional information about the performance of the detector is also available through the PDA Calibration window, accessed from Run Samples in the Empower software.

Verifying detector calibration

Verify the PDA detector calibration after removing and replacing a flow cell. Before you power-on the detector, prime the system to fill the flow cell with solvent and remove any air.

Recommendation: Impurities in the flow cell may affect wavelength verification results. Ensure that the flow cell is clean before you calibrate.

Caution: To ensure the detector is properly aligned and calibrated, the flow cell must be filled with solvent before you power-on the detector. An empty flow cell will cause a calibration error.

To verify detector calibration:

1. Select a solvent compatible with the samples and mobile phases that you have been using. If you have been using buffers, flush with 10 mL of HPLC-grade water, then flush with 10 mL of a low-surface-tension solvent such as methanol.

Tip: Ensure that the solvent is miscible with the previous mobile phase.

- 2. In the console, select 2998 PDA Detector from the system tree.
- 3. In the PDA detector information window, click Maintain > Verify calibration > Start. The test time appears in the Run Time bar graph.
- 4. When the test ends, the Results pane appears. Verify that the detector passed the test.

Rule: The maximum deviation must be within +/-1 nm of the standard calibration to pass.

Tip: If the test fails, flush the flow cell and run the verify detector calibration procedure again.

5. Click Close.

Reading lamp energy

Recommendation: Impurities in the flow cell may affect reading lamp energy. Ensure that the flow cell is clean before you read lamp energy.

To read the lamp energy:

- 1. In the console, select 2998 PDA Detector from the system tree.
- In the PDA detector information window, click Maintain > Read energy > Read. The Read Energy dialog box appears.
- 3. Click Close.

Performing the erbium calibration

Recommendation: Impurities in the flow cell may affect wavelength calibration. Ensure that the flow cell is clean before you perform calibration.

Before you begin this procedure, your detector must be set up and configured as described in Chapter 2.

During the start-up verification sequence, the PDA detector performs an erbium calibration. You can also start this procedure manually.

To start a manual erbium verification:

- 1. In the console, select 2998 PDA Detector from the system tree.
- 2. In the PDA detector information window, click Troubleshoot > Erbium calibration. The Erbium Calibration dialog box appears.
- 3. Click Optimize Exposures.
- 4. Click Start.

Result: The detector moves the filter into the erbium position, locates the 256.5, 378.9, and 521.4 nm erbium absorbance peaks, and locates the 656.1 and 486.1 nm deuterium emission lines.

Rule: The maximum deviation must be within +/-1 nm to pass.

5. Click Stop. The erbium filter moves back to original position.



- This procedure can affect spectral library matching and peak purity calibrations.
- You might need to reacquire spectral libraries each time you perform any calibration procedure. See Chapter 5 for more information.
- 6. Click Save > OK > Close.

Reading the calibration constants

To read the calibration constants:

- 1. In the console, select 2998 PDA Detector from the system tree.
- 2. Click Troubleshoot > Calibration constants. The Calibration Constants dialog box appears.
- 3. Click Read.
- 4. Click Close.

Displaying PDA detector rear panel interface connections

You can use the console to determine the status of input/output signal connections or contact closures on the rear panel of the 2998 PDA detector. This display gives you a real-time status of the instrument's signal connections. A green LED symbol indicates a signal cable is connected to the terminal. A red LED symbol indicates no signal cable is connected to the terminal.

To display 2998 PDA detector rear panel interface connections:

- 1. In the console, select 2998 PDA Detector from the system tree.
- 2. In the PDA detector information window, click Troubleshoot > Rear panel. The PDA Detector Rear Panel dialog box appears.

The following table describes the PDA detector I/O connections.

Signal connections	Description	
Analog 1 (Out)	Method programmable analog output.	
Analog 2 (Out)	Method programmable analog output.	
Switch 1 (Out)	Controls a timed event or threshold level and is a user-programmable auxiliary output.	
Switch 2 (Out)	Controls a timed event or threshold level and is a user-programmable auxiliary output.	
Inject Start (In)	Activates timed events by triggering the run-time clock to start.	
Lamp Off (In)	When the input is triggered, the lamp is turned off.	
Chart Mark (In)	Adds a chart mark (at 10% of full scale) to either or both analog output channels (Signal Out 1 and Signal Out 2) and is configurable.	
Auto Zero (In)	Calculates an offset value that, when added to the sample signal, makes the resulting baseline signal zero.	

PDA detector analog-out/event-in connections

Changing the rear panel interface connections

Via the rear panel display, you can open and close certain output connections. Opening or closing output connections can be useful when you need to start or stop an injection or troubleshoot system connectivity.

To change PDA detector rear panel interface connections:

- 1. In the console, select 2998 PDA Detector from the system tree.
- 2. In the PDA detector information window, click Troubleshoot > Rear Panel.
- 3. In the PDA Detector Rear Panel dialog box, locate a signal connection with an Out label, and then click the red or green LED symbol. The output signal for the selected signal connection indicates whether it is opened] or closed].

General troubleshooting

This section suggests possible causes of errors and recommends troubleshooting actions. Keep in mind that the source of apparent detector problems can actually be the chromatography, or it can involve other system components.

Most detector problems are relatively easy to correct. If, after running the diagnostic functions applicable to your problem and troubleshooting the detector, you cannot correct an error condition, contact Waters' Technical Service department.

Power surges

Power surges, line spikes, and transient energy sources can adversely affect detector operations. Be sure that the electrical supply used for the detector is properly grounded and free from any of these conditions.

Power LED

The power LED, on the detector's front panel, indicates when the instrument is powered-on or powered-off. It is constant green when the instrument is working properly.

Lamp LED

The lamp LED, to the left of the power LED, indicates the lamp status.

LED mode and color	Description
Unlit	Indicates the detector lamp is extinguished.
Constant green	Indicates the detector lamp is ignited and the detector has passed power-up wavelength verification.
Flashing green	Indicates the detector is initializing or verifying wavelength calibration.

Lamp LED indications

Lamp LED indications (Continued)

LED mode and color	Description
Flashing red	Indicates an error stopped the detector. Information regarding the error that caused the failure can be found in the console log. See also: 2998 PDA Detector online Help
Constant red	Indicates a detector failure that prevents further operation. Power-off the detector, and then power-on. If the LED is still steady red, contact your Waters service representative.

Clearing bubbles from the flow cell

To clear bubbles from the flow cell:

Establish liquid flow through the detector flow cell with degassed acetonitrile or methanol at a flow rate that is anticipated to be used for subsequent analysis.

Detector troubleshooting

The following table contains general hardware troubleshooting for the detector.

PDA detector troubleshooting

Symptom	Possible cause	Corrective action
Both LEDs unlit	No power	 Inspect line cord connections. Test electrical outlet for power.
	Open (spent) or defective fuse	Replace fuse (see page 3-10.

Symptom	Possible cause	Corrective action
Change in reference spectrum	Mobile phase contains gas or is contaminated	Prepare fresh mobile phase and degas thoroughly.
	Air bubbles trapped in flow cell	Reseat and check alignment of flow cell. Flush the flow cell (see page 3-4), or apply slight backpressure of 207 to 345 kPa (2 to 3 bar, 30 to 50 psi) on the detector waste outlet. For example, connect a 1- to 2-foot (30- to 60-cm) length of 0.009-inch (0.23-mm) ID tubing connected to the detector waste outlet.
Continuous audible tone	Detector failure	Power the PDA detector off and on again.
Detector not responding to console	Bad or disconnected cable	Inspect cable connections, tighten connectors, or replace cable.
	Configuration problem	Check Ethernet configuration. For details, see Empower online Help.
Lamp light flashes green and power light constant green	The detector is initializing.	No corrective action required. Wait until initialization completes (both lights constant green).

PDA detector troubleshooting (Continued)

Symptom	Possible cause	Corrective action
Lamp light flashes red and power light constant green	Failed startup diagnostic tests	Reseat and check alignment of flow cell (see page 3-5).
		Flush the flow cell (see page 3-4).
	Dirty flow cell causing shutter diagnostic test to fail	Flush the flow cell (see page 3-4).
	Insufficient energy reaching photodiode array because of air bubble	Flush the flow cell (see page 3-4), or apply slight backpressure of 207 to 345 kPa (2 to 3 bar, 30 to 50 psi) on the detector waste outlet. For example, connect a 1- to 2-foot (30- to 60-cm) length of 0.009-inch (0.23-mm) ID tubing connected to the detector waste outlet.
	Weak lamp	Replace the lamp (see page 3-7).
Shutter failure message	Shutter failed	 Clear bubbles from the flow cell (see page 4-7). Power the PDA detector off and on again.

PDA detector troubleshooting (Continued)

Symptom	Possible cause	Corrective action
Solvent in drain line	Leak from flow cell gasket	Replace the flow cell (see page 3-5).
	Leak from flow cell inlet and outlet fittings	Check fittings for overtightening or undertightening, and replace fittings if necessary.

PDA detector troubleshooting (Continued)
5 Spectral Contrast Theory

The spectral contrast algorithm compares the UV/Vis absorbance spectra of samples the detector collects. This chapter describes the theory on which the algorithm is based, explaining how it exploits differences in the shapes of the absorbance spectra. It also explains how spectral contrast represents those spectra as vectors, determining whether differences among them arise from the presence of multiple compounds in the same peaks (coelution) or from nonideal conditions like noise, photometric error, or solvent effects.

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Comparing absorbance spectra

When measured at specific solvent and pH conditions, the shape of a compound's absorbance spectrum characterizes the compound. The varying extent of UV/Vis absorbance occurring at different wavelengths produces a unique spectral shape.

The following figure shows the absorbance spectra for two compounds, A and B. The ratio of the absorbance at 245 nm to that at 257 nm is about 2.2 for compound A and 0.7 for compound B. Note that this comparison of a single wavelength pair's absorbance ratios yields little information about a compound. For more information, you must compare the ratios of multiple wavelength pairs.



Comparing spectra of two compounds

Representing spectra as vectors

The spectral contrast algorithm uses vectors to quantify differences in the shapes of spectra, converting baseline-corrected spectra to vectors and then comparing the vectors. Spectral vectors have two properties:

- Length Proportional to analyte concentration.
- Direction Determined by the relative absorbance of the analyte at all wavelengths (its absorbance spectrum). Direction is independent of concentration for peaks that are less than 1.0 absorbance units (AU) across the collected wavelength range.

Vector direction contributes to the identification of a compound, because the direction is a function of the absorbance compound's spectrum. The ability of spectral vectors to differentiate compounds depends on the resolution of spectral features. As both wavelength range and spectral resolution increase, the precision of a spectral vector for the resultant spectrum increases. A detector-derived vector can include absorbances in the range of 190 to 800 nm. To enhance spectral sensitivity, set the bench resolution to 1.2 nm.

Tip: Do not include wavelengths where there is no analyte absorbance.

Vectors derived from two wavelengths

The spectral contrast algorithm uses vectors to characterize spectra. To understand the vector principle, consider two vectors, in the figure below, which are based on the spectra depicted in the previous figure.



Plotting vectors for two spectra

In this figure, the axes reflect the absorbance units of the two wavelengths used to calculate the absorbance ratio of the previous figure. The head of the vector for Compound A lies at the intersection of the absorbance values (for Compound A), at the two wavelengths represented by each axis. The remaining vector is similarly derived from the spectrum of Compound B.

Compound B's vector points in a direction different from Compound A's. Expressed by the spectral contrast angle (θ), this difference reflects the difference between the two compounds' absorbance ratios at wavelengths 245 nm and 257 nm. A spectral contrast angle greater than zero indicates a shape difference between spectra (see "Spectral contrast angles" on page 5-4).

Finally, note that the length of the vectors is proportional to concentration.

Vectors derived from multiple wavelengths

When absorbance ratios are limited to two wavelengths, the chance that two different spectra share the same absorbance ratio is greater than if comparison is made using absorbance ratios at many wavelengths. Therefore, the spectral contrast algorithm uses absorbances from multiple wavelengths to form a vector in an n-dimensional vector space, where n is the number of wavelengths from the spectrum.

To compare two spectra, the spectral contrast algorithm forms a vector for each spectrum in an n-dimensional space. The two spectral vectors are compared mathematically to compute the spectral contrast angle.

As with the two-wavelength comparison, a spectral contrast angle of zero in *n*-dimensional space means that all ratios of absorbances at corresponding wavelengths match. Conversely, if any comparison of ratios does not match, the corresponding vectors point in different directions.

Spectral contrast angles

Spectra of identical shape have vectors that point in the same direction. Spectra of varying shapes have vectors that point in different directions. The angle between the two vectors of any two spectra, the spectral contrast angle, expresses the difference in direction between the spectral vectors of two spectra.

A spectral contrast angle can vary from 0° to 90° . A spectral contrast angle approaching 0° indicates little shape difference between the compared spectra. Matching a spectrum to itself produces a spectral contrast angle of exactly 0° . The maximum spectral contrast angle, 90° , indicates that the two spectra do not overlap at any wavelength.

To illustrate the relationship between the spectral contrast angle and spectral shape differences, consider the pairs of spectra shown in the next three figures.

Spectra with different shapes

In the following figure, the absorbance spectra of two compounds, A and B, are distinctly different. They therefore produce a large spectral contrast angle (62.3°) .

Spectra that produce a large spectral contrast angle



Spectra with similar shapes

In the following figure, the absorbance spectra of two compounds, A and B, are similar. They therefore produce a small spectral contrast angle (3.0°) .



Spectra with a small spectral contrast angle

Differences between spectra of the same compound

Small but significant differences between absorbance spectra can result from factors other than the absorbance properties of different compounds. For example, multiple spectra of the *same* compound may exhibit slight differences because of detector noise, photometric error, high sample concentration, or variations in solvent conditions. The spectra in the next figure, for example, show how instrument noise can affect the shape of an absorbance spectrum of one compound at two concentrations, one high and one low. Note that the spectral contrast angle between these absorbance spectra of the same compound is 3.4° .



Normalized absorbance spectra of a compound at two concentrations

Undesirable effects

Shape differences between absorbance spectra can be caused by one or more of the following undesirable effects:

- Detector noise
- Photometric error caused by high sample concentration
- Variation in solvent composition

These sources of spectral variation can cause chemically pure, baseline-resolved peaks to exhibit a small level of spectral inhomogeneity. You can assess the significance of spectral inhomogeneity by comparing a spectral contrast angle to a threshold angle (see page 5-8).

Detector noise

Statistical and thermal variations add electronic noise to the detector's absorbance measurements. The noise, which manifests itself as fluctuations in the baseline, is known as baseline noise. The magnitude of any absorbance differences caused by statistical and thermal variations can be predicted from the instrument noise in the baseline region of a chromatogram.

Photometric error

At high absorbances (generally those greater than 1 AU), a combination of effects can produce slight departures (about 1%) from Beer's law because of photometric error. Although photometric errors at this level can negligibly affect quantitation, they can nevertheless be a significant source of spectral inhomogeneity. To minimize the effects of photometric error for all spectral contrast operations, the maximum spectral absorbance of a compound should be less than 1 AU. Keep in mind that the absorbance of the mobile phase reduces the working linear dynamic range by the amount of mobile phase absorbance at each wavelength. For examples of mobile phase absorbance, see Appendix C.

See also: For more information about the effects of the photometric error curve, refer to *Principles of Instrumental Analysis*, 3rd. ed., by Douglas A. Skoog, Saunders College Publishing, 1985, pp. 168–172.

Solvent changes

As long as solvent concentration and composition do not change (isocratic operation), background absorbance, if any, by the solvent remains constant. However, change in solvent pH or composition, such as that which occurs in gradient operation, can affect the intrinsic spectral shape of a compound. (See the figure "Effects of pH on the absorbance spectrum of p-aminobenzoic acid" on page 5-9).

Threshold angle

In addition to computing spectral contrast angles, the spectral contrast algorithm also computes a threshold angle. The threshold angle is the maximum spectral contrast angle between spectra that can be attributed to nonideal phenomena. Comparison of a spectral contrast angle to its threshold angle can help determine whether the shape difference between spectra is significant. In general, a spectral contrast angle less than its threshold angle indicates that shape differences are attributable to nonideal phenomena alone and that no evidence exists for significant differences between the spectra. A spectral contrast angle greater than its threshold angle indicates that the shape differences arise from significant differences between the spectra. When automating the spectral contrast comparison, the maximum absorbance of the spectra must not exceed 1 AU.





A Safety Advisories

Waters instruments display hazard symbols designed to alert you to the hidden dangers of operating and maintaining the instruments. Their corresponding user guides also include the hazard symbols, with accompanying text statements describing the hazards and telling you how to avoid them. This appendix presents all the safety symbols and statements that apply to the entire line of Waters products.

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Warning symbols

Warning symbols alert you to the risk of death, injury, or seriously adverse physiological reactions associated with an instrument's use or misuse. Heed all warnings when you install, repair, and operate Waters instruments. Waters assumes no liability for the failure of those who install, repair, or operate its instruments to comply with any safety precaution.

Task-specific hazard warnings

The following warning symbols alert you to risks that can arise when you operate or maintain an instrument or instrument component. Such risks include burn injuries, electric shocks, ultraviolet radiation exposures, and others.

When the following symbols appear in a manual's narratives or procedures, their accompanying text identifies the specific risk and explains how to avoid it.

Warning: (General risk of danger. When this symbol appears on an instrument, consult the instrument's user documentation for important safety-related information before you use the instrument.)

Warning: (Risk of burn injury from contacting hot surfaces.)



Warning: (Risk of fire.)

Warning: (Risk of sharp-point puncture injury.)

Warning: (Risk of hand crush injury.)

Warning: (Risk of exposure to ultraviolet radiation.)



Warning: (Risk of contacting corrosive substances.)

Warning: (Risk of exposure to a toxic substance.)

Warning: (Risk of personal exposure to laser radiation.)

Warning: (Risk of exposure to biological agents that can pose a serious health threat.)



Specific warnings

The following warnings can appear in the user manuals of particular instruments and on labels affixed to them or their component parts.

Burst warning

This warning applies to Waters instruments fitted with nonmetallic tubing.



Warning: Pressurized nonmetallic, or polymer, tubing can burst. Observe these precautions when working around such tubing:

- Wear eye protection.
- Extinguish all nearby flames.
- Do not use tubing that is, or has been, stressed or kinked.
- Do not expose nonmetallic tubing to incompatible compounds like tetrahydrofuran (THF) and nitric or sulfuric acids.
- Be aware that some compounds, like methylene chloride and dimethyl sulfoxide, can cause nonmetallic tubing to swell, which significantly reduces the pressure at which the tubing can rupture.

Mass spectrometer flammable solvents warning

This warning applies to instruments operated with flammable solvents.

Warning: Where significant quantities of flammable solvents are involved, a continuous flow of nitrogen into the ion source is required to prevent possible ignition in that enclosed space.

Ensure that the nitrogen supply pressure never falls below 690 kPa (6.9 bar, 100 psi) during an analysis in which flammable solvents are used. Also ensure a gas-fail connection is connected to the LC system so that the LC solvent flow stops if the nitrogen supply fails.

Mass spectrometer shock hazard

This warning applies to all Waters mass spectrometers.

Warning: To avoid electric shock, do not remove the mass spectrometer's protective panels. The components they cover are not user-serviceable.

This warning applies to certain instruments when they are in Operate mode.

Warning: High voltages can be present at certain external surfaces of the mass spectrometer when the instrument is in Operate mode. To avoid non-lethal electric shock, make sure the instrument is in Standby mode before touching areas marked with this high voltage warning symbol.

Biohazard warning

This warning applies to Waters instruments that can be used to process material that might contain biohazards: substances that contain biological agents capable of producing harmful effects in humans.

Warning: Waters instruments and software can be used to analyze or process potentially infectious human-sourced products, inactivated microorganisms, and other biological materials. To avoid infection with these agents, assume that all biological fluids are infectious, observe Good Laboratory Practices, and consult your organization's biohazard safety representative regarding their proper use and handling. Specific precautions appear in the latest edition of the US National Institutes of Health (NIH) publication, *Biosafety in Microbiological and Biomedical Laboratories* (BMBL).

Chemical hazard warning

This warning applies to Waters instruments that can process corrosive, toxic, flammable, or other types of hazardous material.

Warning: Waters instruments can be used to analyze or process potentially hazardous substances. To avoid injury with any of these materials, familiarize yourself with the materials and their hazards, observe Good Laboratory Practices (GLP), and consult your organization's safety representative regarding proper use and handling. Guidelines are provided in the latest edition of the National Research Council's publication, *Prudent Practices in the Laboratory: Handling and Disposal of Chemicals.*

Caution symbol

The caution symbol signifies that an instrument's use or misuse can damage the instrument or compromise a sample's integrity. The following symbol and its associated statement are typical of the kind that alert you to the risk of damaging the instrument or sample.



Caution: To avoid damage, do not use abrasives or solvents to clean the instrument's case.

Warnings that apply to all Waters instruments

When operating this device, follow standard quality control procedures and the equipment guidelines in this section.

Attention: Changes or modifications to this unit not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment.

Important: Toute modification sur cette unité n'ayant pas été expressément approuvée par l'autorité responsable de la conformité à la réglementation peut annuler le droit de l'utilisateur à exploiter l'équipement.

Achtung: Jedwede Änderungen oder Modifikationen an dem Gerät ohne die ausdrückliche Genehmigung der für die ordnungsgemäße Funktionstüchtigkeit verantwortlichen Personen kann zum Entzug der Bedienungsbefugnis des Systems führen.

Avvertenza: qualsiasi modifica o alterazione apportata a questa unità e non espressamente autorizzata dai responsabili per la conformità fa decadere il diritto all'utilizzo dell'apparecchiatura da parte dell'utente.

Atencion: cualquier cambio o modificación efectuado en esta unidad que no haya sido expresamente aprobado por la parte responsable del cumplimiento puede anular la autorización del usuario para utilizar el equipo.

注意:未經有關法規認證部門允許對本設備進行的改變或修改,可能會使使用者喪失操作該設備的權利。

注意: 未经有关法规认证部门明确允许对本设备进行的改变或改装,可能会使使用者丧失操 作该设备的合法性。

주의: 규정 준수를 책임지는 당사자의 명백한 승인 없이 이 장치를 개조 또는 변경할 경우, 이 장치를 운용할 수 있는 사용자 권한의 효력을 상실할 수 있습니다.

注意:規制機関から明確な承認を受けずに本装置の変更や改造を行うと、本装置のユー ザーとしての承認が無効になる可能性があります。



Warning: Use caution when working with any polymer tubing under pressure:

- Always wear eye protection when near pressurized polymer tubing.
 - Extinguish all nearby flames.
- Do not use tubing that has been severely stressed or kinked.
- Do not use nonmetallic tubing with tetrahydrofuran (THF) or concentrated nitric or sulfuric acids.
- Be aware that methylene chloride and dimethyl sulfoxide cause nonmetallic tubing to swell, which greatly reduces the rupture pressure of the tubing.

Attention: Manipulez les tubes en polymère sous pression avec precaution:

- Portez systématiquement des lunettes de protection lorsque vous vous trouvez à proximité de tubes en polymère pressurisés.
- Eteignez toute flamme se trouvant à proximité de l'instrument.
- Evitez d'utiliser des tubes sévèrement déformés ou endommagés.
- Evitez d'utiliser des tubes non métalliques avec du tétrahydrofurane (THF) ou de l'acide sulfurique ou nitrique concentré.
- Sachez que le chlorure de méthylène et le diméthylesulfoxyde entraînent le gonflement des tuyaux non métalliques, ce qui réduit considérablement leur pression de rupture.

Vorsicht: Bei der Arbeit mit Polymerschläuchen unter Druck ist besondere Vorsicht angebracht:

- In der Nähe von unter Druck stehenden Polymerschläuchen stets Schutzbrille tragen.
- · Alle offenen Flammen in der Nähe löschen.
- Keine Schläuche verwenden, die stark geknickt oder überbeansprucht sind.
- Nichtmetallische Schläuche nicht für Tetrahydrofuran (THF) oder konzentrierte Salpeter- oder Schwefelsäure verwenden.
- Durch Methylenchlorid und Dimethylsulfoxid können nichtmetallische Schläuche quellen; dadurch wird der Berstdruck des Schlauches erheblich reduziert.

Attenzione: fare attenzione quando si utilizzano tubi in materiale polimerico sotto pressione:

- Indossare sempre occhiali da lavoro protettivi nei pressi di tubi di polimero pressurizzati.
- Spegnere tutte le fiamme vive nell'ambiente circostante.
- Non utilizzare tubi eccessivamente logorati o piegati.
- Non utilizzare tubi non metallici con tetraidrofurano (THF) o acido solforico o nitrico concentrati.
- Tenere presente che il cloruro di metilene e il dimetilsolfossido provocano rigonfiamenti nei tubi non metallici, riducendo notevolmente la pressione di rottura dei tubi stessi.

Advertencia: se recomienda precaución cuando se trabaje con tubos de polímero sometidos a presión:

- El usuario deberá protegerse siempre los ojos cuando trabaje cerca de tubos de polímero sometidos a presión.
- Si hubiera alguna llama las proximidades.
- No se debe trabajar con tubos que se hayan doblado o sometido a altas presiones.
- Es necesario utilizar tubos de metal cuando se trabaje con tetrahidrofurano (THF) o ácidos nítrico o sulfúrico concentrados.
- Hay que tener en cuenta que el cloruro de metileno y el sulfóxido de dimetilo dilatan los tubos no metálicos, lo que reduce la presión de ruptura de los tubos.

警告: 當在有壓力的情況下使用聚合物管線時, 小心注意以下幾點。

- 當接近有壓力的聚合物管線時一定要戴防護眼鏡。
- 熄滅附近所有的火焰。
- 不要使用已經被壓癟或嚴重彎曲管線。
- 不要在非金屬管線中使用四氫呋喃或濃硝酸或濃硫酸。
- 要了解使用二氯甲烷及二甲基亞楓會導致非金屬管線膨脹,大大降低管線的耐壓能力。

▲ 警告:当有压力的情况下使用管线时,小心注意以下几点:

- 当接近有压力的聚合物管线时一定要戴防护眼镜。
 - 熄灭附近所有的火焰。
 - 不要使用已经被压瘪或严重弯曲的管线。
 - 不要在非金属管线中使用四氢呋喃或浓硝酸或浓硫酸。
 - 要了解使用二氯甲烷及二甲基亚枫会导致非金属管线膨胀,大大降低管线的耐压能力。

경고: 가압 폴리머 튜브로 작업할 경우에는 주의하십시오.

- 가압 폴리머 튜브 근처에서는 항상 보호 안경을 착용하십시오.
- 근처의 화기를 모두 끄십시오.
- 심하게 변형되거나 꼬인 튜브는 사용하지 마십시오.
- 비금속(Nonmetallic) 튜브를 테트라히드로푸란(Tetrahydrofuran: THF) 또는 농축 질산 또는 황산과 함께 사용하지 마십시오.
- 염화 메틸렌(Methylene chloride) 및 디메틸술폭시드(Dimethyl sulfoxide)는 비금속 튜브를 부풀려 튜브의 파열 압력을 크게 감소시킬 수 있으므로 유의하십시오.

警告: 圧力のかかったポリマーチューブを扱うときは、注意してください。

- ・ 加圧されたポリマーチューブの付近では、必ず保護メガネを着用してください。
- 近くにある火を消してください。
- 著しく変形した、または折れ曲がったチューブは使用しないでください。
- ・ 非金属チューブには、テトラヒドロフラン(THF)や高濃度の硝酸または硫酸などを流 さないでください。
- 塩化メチレンやジメチルスルホキシドは、非金属チューブの膨張を引き起こす場合があり、その場合、チューブは極めて低い圧力で破裂します。

Warning: The user shall be made aware that if the equipment is used in a manner not specified by the manufacturer, the protection provided by the equipment may be impaired.

Attention: L'utilisateur doit être informé que si le matériel est utilisé d'une façon non spécifiée par le fabricant, la protection assurée par le matériel risque d'être défectueuses.

Vorsicht: Der Benutzer wird darauf aufmerksam gemacht, dass bei unsachgemäßer Verwenddung des Gerätes die eingebauten Sicherheitseinrichtungen unter Umständen nicht ordnungsgemäß funktionieren.

Attenzione: si rende noto all'utente che l'eventuale utilizzo dell'apparecchiatura secondo modalità non previste dal produttore può compromettere la protezione offerta dall'apparecchiatura.

Advertencia: el usuario deberá saber que si el equipo se utiliza de forma distinta a la especificada por el fabricante, las medidas de protección del equipo podrían ser insuficientes.

警告: 使用者必须非常清楚如果設備不是按照製造廠商指定的方式使用, 那麼該設備所提供的保護將被消弱。

警告: 使用者必须非常清楚如果设备不是按照制造厂商指定的方式使用, 那么该设备所提供的保护将被削弱。

경고: 제조업체가 명시하지 않은 방식으로 장비를 사용할 경우 장비가 제공하는 보호 수단이 제대로 작동하지 않을 수 있다는 점을 사용자에게 반드시 인식시켜야 합니다.

警告: ユーザーは、製造元により指定されていない方法で機器を使用すると、機器が提供している保証が無効になる可能性があることに注意して下さい。

Warning: To protect against fire, replace fuses with those of the type and rating printed on panels adjacent to instrument fuse covers.
Attention: pour éviter tout risque d'incendie, remplacez toujours les fusibles par d'autres du type et de la puissance indiqués sur le panneau à proximité du couvercle de la boite à fusible de l'instrument.
Vorsicht: Zum Schutz gegen Feuer die Sicherungen nur mit Sicherungen ersetzen, deren Typ und Nennwert auf den Tafeln neben den Sicherungsabdeckungen des Geräts gedruckt sind.
Attenzione: per garantire protezione contro gli incendi, sostituire i fusibili con altri dello stesso tipo aventi le caratteristiche indicate sui pannelli adiacenti alla copertura fusibili dello strumento.
Advertencia: Para evitar incendios, sustituir los fusibles por aquellos del tipo y características impresos en los paneles adyacentes a las cubiertas de los fusibles del instrumento.

警告:為了避免火災,更換保險絲時,請使用與儀器保險絲蓋旁面板上所印刷之相同類型與規格的保險絲。

警告:为了避免火灾,应更换与仪器保险丝盖旁边面板上印刷的类型和规格相同的保险丝。



경고: 화재의 위험을 막으려면 기기 퓨즈 커버에 가까운 패널에 인쇄된 것과 동일한 타입 및 정격의 제품으로 퓨즈를 교체하십시오.

警告:火災予防のために、ヒューズ交換では機器ヒューズカバー脇のパネルに記載されているタイプおよび定格のヒューズをご使用ください。

Electrical symbols

These can appear in instrument user manuals and on the instrument's front or rear panels.

	Electrical power on
0	Electrical power off
\bigcirc	Standby
	Direct current
\sim	Alternating current
	Protective conductor terminal
m	Frame, or chassis, terminal
	Fuse
X	Recycle symbol: Do not dispose in municipal waste.

Handling symbols

These handling symbols and their associated text can appear on labels affixed to the outer packaging of Waters instrument and component shipments.

<u> 11</u>	Keep upright!
×	Keep dry!
Y	Fragile!
\mathbf{X}	Use no hooks!

A-14 Safety Advisories

B Specifications

This appendix contains product specifications for the 2998 PDA detector.

Attribute	Specification
Height	19.4 cm (7.64 inches)
Depth	61 cm (24.0 inches)
Width	34.3 cm (13.5 inches)
Weight	12.6 kg (27.75 pounds)

Physical specifications

Environmental specifications

Attribute	Specification
Operating temperature	4 to 40 °C (39.2 to 104 °F)
Operating humidity	<95%, noncondensing
Shipping and storage temperature	-30 to 60 °C (-22 to 140 °F)
Shipping and storage humidity	20 to 85%, noncondensing
Acoustic noise (instrument generated)	<50 dBA

Electrical specifications

Attribute	Specification
Protection class ^a	Class I
Overvoltage category ^b	II
Pollution degree ^c	2

Electrical specifications (Continued)

Attribute	Specification
Moisture protection ^d	Normal (IPXO)
Line voltages, nominal	Grounded AC
Voltage range	100 to 240 VAC nominal
Frequency	50 to 60 Hz
Fuse	Two fuses, 100 to 240 VAC, 50 to 60-Hz, F 3.15-A, 250-V (fast-blow), 5 × 20 mm (IEC)
Power consumption	185 VA nominal

- a. **Protection Class I** The insulating scheme used in the instrument to protect from electrical shock. Class I identifies a single level of insulation between live parts (wires) and exposed conductive parts (metal panels), in which the exposed conductive parts are connected to a grounding system. In turn, this grounding system is connected to the third pin (ground pin) on the electrical power cord plug.
- b. **Overvoltage Category II** Pertains to instruments that receive their electrical power from a local level such as an electrical wall outlet.
- c. **Pollution Degree 2** A measure of pollution on electrical circuits, which may produce a reduction of dielectric strength or surface resistivity. Degree 2 refers only to normally nonconductive pollution. Occasionally, however, expect a temporary conductivity caused by condensation.
- d. **Moisture Protection** Normal (IPXO) IPXO means that *no* Ingress Protection against any type of dripping or sprayed water exists. The X is a placeholder that identifies protection against dust, if applicable.

Attribute	Specification
Wavelength range	190 to 800 nm
Optical resolution	1.2 nm
Photodiodes	512
Digital resolution	1.2 nm/pixel
Wavelength accuracy	±1.0 nm
Wavelength repeatability	±0.1 nm
Digital filter	Variable with data rate
Second-order filter	Fixed 371 nm to 800 nm

Performance specifications

Attribute	Specification			
Noise – UV (dry)	10 μAU peak to peak			
	Filter = 1 second, 30-second segments			
	Wavelength = 254 nm			
	Bandwidth = 3.6 nm (3-pixel bunch)			
	Flow cell = analytical, 10 mm			
	Data rate = 2 Hz			
Noise – UV (wet ^a)	10 µAU peak to peak			
	Filter = 1 second, 30-second segments			
	Wavelength = 254 nm			
	Bandwidth = 3.6 nm (3-pixel bunch)			
	Flow cell = analytical, 10 mm			
	Data rate = 2 Hz			
	Flow rate = 0.5 mL/min			
	Solvent = water/acetonitrile, 90/10			
Drift – UV (dry)	1000 µAU/hr			
	Filter = 1 second, 30-second segments			
	Wavelength = 254 nm			
	Bandwidth = 3.6 nm (3-pixel bunch)			
	Warm-up time = 60 minutes			
	Environmental stability = ± 2 °C/hr			
	Flow cell = analytical, 10 mm			
	Data rate = 2 Hz			
Linearity	< 5% at 2.0 AU, propylparaben series at 257 nm, analytical flowcell			
Data rate	1, 2, 5, 10, 20, 40, and 80			

Performance specifications (Continued)

a. Wet testing should be done with water/acetonitrile, 90/10, to minimize the effects of oxygen at 230 nm. Water/methanol, 90/10, can be substituted with appropriate solvent conditioning.

	Waters	2998	PDA	flow	cell	specifications
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Description	Volume (µL)	Pathlength (mm)	Tubing internal diameter (inches)		Pressure rating (bar/psi)	
			Inlet	Outlet	(bai/psi)	
Analytical	9.3	10	0.010	0.010	69/1000	
Auto-purification						
 analytical 	12.4	0.5	0.010	0.040	138/2000	
• prep			0.040			
Microbore	4.1	8.0	0.005	0.005	69/1000	
Semi-preparative	18.3	3.0	0.020	0.020	69/1000	
Cuvette	N/A	10.0	N/A	N/A	N/A	

C Solvent Considerations

This appendix contains certain solvent considerations you must take into account when operating or maintaining the 2998 PDA Detector.

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Warning: To avoid chemical hazards, always observe Good Laboratory Practices when operating your system.

Introduction

Clean solvents

Clean solvents provide reproducible results and permit you to operate with minimal instrument maintenance.

A dirty solvent can cause baseline noise and drift. It can also block the solvent filters with particulate matter.

Solvent quality

Use HPLC-grade solvents for the best possible results. Filter solvents through 0.45-µm filters before their use. Solvents distilled in glass generally maintain their purity from lot to lot; use them to ensure the best possible results.

Preparation checklist

The following solvent preparation guidelines help to ensure stable baselines and good resolution:

- Filter solvents with a 0.45- μm filter.
- Degas and/or sparge the solvent.
- Stir the solvent.
- Keep in a place free from drafts and shock.

Water

Use water only from a high-quality water purification system. If the water system does not deliver filtered water, filter it through a 0.45-µm membrane filter before use.

Using buffers

When you use buffers, dissolve salts first, adjust the pH, then filter to remove insoluble material.

Tetrahydrofuran

When using unstabilized tetrahydrofuran, ensure that your solvent is fresh. Previously opened bottles of tetrahydrofuran contain peroxide contaminants, which cause baseline drift.



Warning: Tetrahydrofuran contaminants (peroxides) are potentially explosive if concentrated or taken to dryness.

Solvent miscibility

Before you change solvents, refer to the table below to determine the miscibility of the solvents to be used. When you change solvents, be aware that

- changes involving two miscible solvents may be made directly. Changes involving two solvents that are not totally miscible (for example, from chloroform to water), require an intermediate solvent (such as isopropanol).
- temperature affects solvent miscibility. If you are running a high-temperature application, consider the effect of the higher temperature on solvent solubility.
- buffers dissolved in water can precipitate when mixed with organic solvents.

When you switch from a strong buffer to an organic solvent, flush the buffer out of the system with distilled water before you add the organic solvent.

Polarity Index	Solvent	Viscosity CP, 20 °C	Boiling Point °C (1 atm)	Miscibility Number (M)	λ Cutoff (nm)
-0.3	N-decane	0.92	174.1	29	—
-0.4	Iso-octane	0.50	99.2	29	210
0.0	N-hexane	0.313	68.7	29	—
0.0	Cyclohexane	0.98	80.7	28	210
1.7	Butyl ether	0.70	142.2	26	
1.8	Triethylamine	0.38	89.5	26	

Solvent miscibility

Polarity Index	Solvent	Viscosity CP, 20 °C	Boiling Point °C (1 atm)	Miscibility Number (M)	λ Cutoff (nm)
2.2	Isopropyl ether	0.33	68.3	—	220
2.3	Toluene	0.59	100.6	23	285
2.4	P-xylene	0.70	138.0	24	290
3.0	Benzene	0.65	80.1	21	280
3.3	Benzyl ether	5.33	288.3	—	
3.4	Methylene chloride	0.44	39.8	20	245
3.7	Ethylene chloride	0.79	83.5	20	
3.9	Butyl alcohol	3.00	117.7	- _	
3.9	Butanol	3.01	177.7	15	
4.2	Tetrahydrofuran	0.55	66.0	17	220
4.3	Ethyl acetate	0.47	77.1	19	260
4.3	1-propanol	2.30	97.2	15	210
4.3	2-propanol	2.35	117.7	15	_
4.4	Methyl acetate	0.45	56.3	15, 17	260
4.5	Methyl ethyl ketone	0.43	80.0	17	330
4.5	Cyclohexanone	2.24	155.7	28	210
4.5	Nitrobenzene	2.03	210.8	14, 20	—
4.6	Benzonitrile	1.22	191.1	15, 19	—
4.8	Dioxane	1.54	101.3	17	220
5.2	Ethanol	1.20	78.3	14	210
5.3	Pyridine	0.94	115.3	16	305
5.3	Nitroethane	0.68	114.0	—	—
5.4	Acetone	0.32	56.3	15, 17	330
5.5	Benzyl alcohol	5.80	205.5	13	
5.7	Methoxyethanol	1.72	124.6	13	
6.2	Acetonitrile	0.37	81.6	11, 17	190
6.2	Acetic acid	1.26	117.9	14	
6.4	Dimethylformamide	0.90	153.0	12	

Solvent miscibility (Continued)

Polarity Index	Solvent	Viscosity CP, 20 °C	Boiling Point °C (1 atm)	Miscibility Number (M)	λ Cutoff (nm)
6.5	Dimethylsulfoxide	2.24	189.0	9	—
6.6	Methanol	0.60	64.7	12	210
7.3	Formamide	3.76	210.5	3	
9.0	Water	1.00	100.0		

Solvent miscibility (Continued)

How to use miscibility numbers

Use miscibility numbers (M-numbers) to predict the miscibility of a liquid with a standard solvent (see page C-3.)

To predict the miscibility of two liquids, subtract the smaller M-number value from the larger M-number value.

- If the difference between the two M-numbers is 15 or less, the two liquids are miscible in all proportions at 15 °C.
- A difference of 16 indicates a critical solution temperature from 25 °C to 75 °C, with 50 °C as the optimal temperature.
- If the difference is 17 or greater, the liquids are immiscible or their critical solution temperature is above 75 °C.

Some solvents prove immiscible with solvents at both ends of the lipophilicity scale. These solvents receive a dual M-number:

- The first number, always lower than 16, indicates the degree of miscibility with highly lipophilic solvents.
- The second number applies to the opposite end of the scale. A large difference between these two numbers indicates a limited range of miscibility.

For example, some fluorocarbons are immiscible with all the standard solvents and have M-numbers of 0, 32. Two liquids with dual M-numbers are usually miscible with each other.

A liquid is classified in the M-number system by testing for miscibility with a sequence of standard solvents. A correction term of 15 units is then either added or subtracted from the cutoff point for miscibility.

Buffered solvents

When using a buffer, use a good quality reagent and filter it through a 0.45-µm filter.

Do not leave the buffer stored in the system after use. Flush all fluid line pathways with HPLC-quality water before shutting the system down and leave distilled water in the system (flush with 90% HPLC-quality water: 10% methanol for shutdowns scheduled to be more than one day). Use a minimum of 15 mL for sparge-equipped units, and a minimum of 45 mL for in-line vacuum degasser-equipped units.

Head height

Position the solvent reservoirs at a level above the HPLC equipment or on top of the pump or detector (with adequate spill protection).

Minimum tubing bend radius recommendations

When bending tubing, consult the following table. The bend radius on your tubing should not be less than the arc shown in the table. The scale is 1:1 so that the diagram can be used as a template.

Tubing size (OD)	Minimum bend radius
1/16-inch or smaller tubing	1/4-inch

Minimum bend radius for stainless steel tubing

Tubing size (OD)	Minimum bend radius
1/8-inch tubing	1/2-inch
	R

Minimum bend radius for stainless steel tubing (Continued)

Solvent viscosity

Generally, viscosity is not important when you are operating with a single solvent or under low pressure. However, when you are running a gradient, the viscosity changes that occur as the solvents are mixed in different proportions can result in pressure changes during the run. For example, a 1:1 mixture of water and methanol produces twice the pressure of either water or methanol alone.

If the extent to which the pressure changes will affect the analysis is not known, monitor the pressure during the run using the Chart Out terminal.

Mobile phase solvent degassing

Mobile phase difficulties account for 70% or more of all liquid chromatographic problems. Using degassed solvents is important, especially at wavelengths below 220 nm.

Degassing provides

- stable baselines and enhanced sensitivity.
- reproducible retention times for eluting peaks.
- reproducible injection volumes for quantitation.
- stable pump operation.

Gas solubility

Only a finite amount of gas can be dissolved in a given volume of liquid. This amount depends on

- the chemical affinity of the gas for the liquid.
- the temperature of the liquid.
- the pressure applied to the liquid.

Changes in the composition, temperature, or pressure of the mobile phase can all lead to outgassing.

Effects of intermolecular forces

Nonpolar gases (N_2 , O_2 , CO_2 , He) are more soluble in nonpolar solvents than in polar solvents. Generally, a gas is most soluble in a solvent with intermolecular attractive forces similar to those in the gas (like dissolves like).

Effects of temperature

Temperature affects the solubility of gases. If the heat of solution is exothermic, the solubility of the gas decreases when you heat the solvent. If the heat of solution is endothermic, the solubility increases when you heat the solvent. For example, the solubility of He in $\rm H_2O$ decreases with an increase in temperature, but the solubility of He in benzene increases with an increase in temperature.

Effects of partial pressure

The mass of gas dissolved in a given volume of solvent is proportional to the partial pressure of the gas in the vapor phase of the solvent. If you decrease the partial pressure of the gas, the amount of that gas in solution also decreases.

Solvent degassing methods

This section describes the solvent degassing techniques that will help you to attain a stable baseline. Degassing your solvent also improves reproducibility and pump performance.

You can use either of the following methods to degas solvents:

- Sparging with helium
- Vacuum degassing
Sparging

Sparging removes gases from solution by displacing dissolved gases in the solvent with a less soluble gas, usually helium. Well-sparged solvent improves pump performance. Helium sparging brings the solvent to a state of equilibrium, which may be maintained by slow sparging or by keeping a blanket of helium over the solvent. Blanketing inhibits reabsorption of atmospheric gases.

Sparging can change the composition of mixed solvents.

Vacuum degassing

The in-line vacuum degasser operates on the principle of Henry's Law to remove dissolved gases from the solvent. Henry's Law states that the mole fraction of a gas dissolved in liquid is proportional to the partial pressure of that gas in the vapor phase above the liquid. If the partial pressure of a gas on the surface of the liquid is reduced, for example, by evacuation, then a proportional amount of that gas comes out of solution.

Vacuum degassing can change the composition of mixed solvents.

Solvent degassing considerations

Select the most efficient degassing operation for your application. To remove dissolved gas quickly, consider the following.

Sparging

Helium sparging gives stable baselines and better sensitivity than sonication in a detector, and prevents reabsorption of atmospheric gases. Use this method to retard oxidation when you are using tetrahydrofuran or other peroxide-forming solvents.

Vacuum Degassing

The longer the solvent is exposed to the vacuum, the more dissolved gases are removed. Two factors affect the amount of time the solvent is exposed to the vacuum:

- Flow rate At low flow rates, most of the dissolved gas is removed as the solvent passes through the vacuum chamber. At higher flow rates, lesser amounts of gas per unit volume of solvent are removed.
- Surface area of the degassing membrane The length of the degassing membrane is fixed in each vacuum chamber. To increase the length of membrane, you can connect two or more vacuum chambers in series.

The in-line degasser is available as an option or factory-installed in the Waters[®] 2695 Separations Module, XE model.

Wavelength selection

This section includes UV cutoff ranges for

- common solvents.
- common mixed mobile phases.
- chromophores.

UV cutoffs for common solvents

The table below shows the UV cutoff (the wavelength at which the absorbance of the solvent is equal to 1 AU) for some common chromatographic solvents. Operating at a wavelength near or below the cutoff increases baseline noise because of the absorbance of the solvent.

Solvent	UV Cutoff (nm)	Solvent	UV Cutoff (nm)
1-Nitropropane	380	Ethylene glycol	210
2-Butoxyethanol	220	Iso-octane	215
Acetone	330	Isopropanol	205
Acetonitrile	190	Isopropyl chloride	225

UV cutoff wavelengths for common chromatographic solvents

Solvent	UV Cutoff (nm)	Solvent	UV Cutoff (nm)
Amyl alcohol	210	Isopropyl ether	220
Amyl chloride	225	Methanol	205
Benzene	280	Methyl acetate	260
Carbon disulfide	380	Methyl ethyl ketone	330
Carbon tetrachloride	265	Methyl isobutyl ketone	334
Chloroform	245	Methylene chloride	233
Cyclohexane	200	<i>n</i> -Pentane	190
Cyclopentane	200	<i>n</i> -Propanol	210
Diethyl amine	275	<i>n</i> -Propyl chloride	225
Dioxane	215	Nitromethane	380
Ethanol	210	Petroleum ether	210
Ethyl acetate	256	Pyridine	330
Ethyl ether	220	Tetrahydrofuran	230
Ethyl sulfide	290	Toluene	285
Ethylene dichloride	230	Xylene	290

UV cutoff wavelengths for common chromatographic solvents (Continued)

Mixed mobile phases

The table below contains approximate wavelength cutoffs for some other solvents, buffers, detergents, and mobile phases. The solvent concentrations represented are those most commonly used. If you want to use a different concentration, you can determine approximate absorbance using Beer's Law, because absorbance is proportional to concentration.

Mobile Phase	UV Cutoff (nm)	Mobile Phase	UV Cutoff (nm)
Acetic acid, 1%	230	Sodium chloride, 1 M	207
Ammonium acetate, 10 mM	205	Sodium citrate, 10 mM	225
Ammonium bicarbonate, 10 mM	190	Sodium dodecyl sulfate	190
BRIJ 35, 0.1%	190	Sodium formate, 10 mM	200
CHAPS, 0.1%	215	Triethyl amine, 1%	235
Diammonium phosphate, 50 mM	205	Trifluoracetic acid, 0.1%	190
EDTA, disodium, 1 mM	190	TRIS HCl, 20 mM, pH 7.0, pH 8.0	202, 212
HEPES, 10 mM, pH 7.6	225	Triton-X [™] 100, 0.1%	240
Hydrochloric acid, 0.1%	190	Waters PIC [®] Reagent A, 1 vial/liter	200
MES, 10 mM, pH 6.0	215	Waters PIC Reagent B-6, 1 vial/liter	225
Potassium phosphate, monobasic, 10 mM dibasic, 10 mM	190 190	Waters PIC Reagent B-6, low UV, 1 vial/liter	190
Sodium acetate, 10 mM	205	Waters PIC Reagent D-4, 1 vial/liter	190

Wavelength cutoffs for different mobile phases

Wavelength selection for chromophore detection

Certain functional groups found in most compounds absorb light selectively. These groups, known as chromophores, and their behavior can be used to categorize the detection of sample molecules.

The table below lists some common chromophores, and their detection wavelengths (λ_{max}), as well as the molar absorptivity (ϵ_{max}) of each group. Use this information as a guide to select the optimal operating wavelength for a particular analysis. Because of the diversity possible within a given sample, scanning over a range of wavelengths may be necessary to determine the best wavelength for a particular analysis.

Chromophore	Chemical Configuration	λ _{max} (nm)	∈ _{max} (L/m/cm)	λ _{max} (nm)	∈ _{max} (L/m/cm)
Ether	-0	185	1000		
Thioether	—S—	194	4600	215	1600
Amine	-NH ₂	195	2800		
Thiol	—SH	195	1400		
Disulfide	SS	194	5500	255	400
Bromide	—Br	208	300		
Iodide	—I	260	400		
Nitrile	—C≡N	160	_		
Acetylide	—C≡C—	175-180	6000		
Sulfone	_SO ₂	180			
Oxime	—NOH	190	5000		
Azido	>C=N—	190	5000		
Ethylene	—C=C—	190	8000		
Ketone	>C=O	195	1000	270-285	18-30
Thioketone	>C=S	205	strong		
Esters	—COOR	205	50		
Aldehyde	—СНО	210	strong	280-300	11-18
Carboxyl	—СООН	200-210	50-70	1	
Sulfoxide	>S—0	210	1500		

Electronic absorption bands of representative chromophores *

Chromophore	Chemical Configuration	λ _{max} (nm)	∈ _{max} (L/m/cm)	λ _{max} (nm)	∈ _{max} (L/m/cm)
Nitro	-NO ₂	210	strong		
Nitrile	-ONO	220-230	1000-2000	300-400	10
Azo	N=N	285-400	3-25		
Nitroso	—N=O	302	100		
Nitrate	-ONO ₂	270 (shoulde r)	12		
Allene	—(C=C) ₂ — (acyclic)	210-230	21,000		
Allene	—(C=C)3—	260	35,000		
Allene	(C=C)4	300	52,000		
Allene	—(C=C)5—	330	118,000		
Allene	(C=C)2 (alicyclic)	230-260	3000-8000		
Ethylenic/ Acetylenic	C=C—C≡C	219	6,500		
Ethylenic/ Amido	C=C-C=N	220	23,000		
Ethylenic/ Carbonyl	C=C—C=O	210-250	10,000- 20,000		
Ethylenic/ Nitro	C=C—NO ₂	229	9,500		

Electronic absorption bands of representative chromophores (Continued)*

*Willard, H. H. and others. *Instrumental Methods of Analysis*, 6th ed. Litton Educational Publishing, Inc., 1981. Reprinted by permission of Wadsworth Publishing Co., Belmont, California, 94002.

Mobile phase absorbance

This section lists the absorbances at several wavelengths for frequently used mobile phases. Choose the mobile phase carefully to reduce baseline noise.

The best mobile phase for your application is one that is transparent at the chosen detection wavelengths. With such a mobile phase, ensure that any

absorbance is due only to the sample. Absorbance by the mobile phase also reduces the linear dynamic range of the detector by the amount of absorbance the autozero function cancels, or "autozeroes," out. Wavelength, pH, and concentration of the mobile phase affects its absorbance. Examples of several mobile phases are given in the table below.

Tip: The absorbances in the table below are based on a 10-mm pathlength.

	Absor	Absorbance at specified wavelength (nm)									
	200	205	210	215	220	230	240	250	260	280	
Solvents											
Acetonitrile	0.05	0.03	0.02	0.01	0.01	< 0.01			_	—	
Methanol (not degassed)	2.06	1.00	0.53	0.37	0.24	0.11	0.05	0.02	< 0.01		
Methanol (degassed)	1.91	0.76	0.35	0.21	0.15	0.06	0.02	< 0.01			
Isopropanol	1.80	0.68	0.34	0.24	0.19	0.08	0.04	0.03	0.02	0.02	
Unstablized tetrahydro- furan (THF), fresh	2.44	2.57	2.31	1.80	1.54	0.94	0.42	0.21	0.09	0.05	
Unstablized tetrahydro- furan (THF), old	>2.5	>2.5	>2.5	>2.5	>2.5	>2.5	>2.5	>2.5	2.5	1.45	

Mobile phase absorbance measured against air or water

Mobile phase absorbance measured against air or water (Continued)

	Absorbance at specified wavelength (nm)										
	200	205	210	215	220	230	240	250	260	280	
Acids and bases											
Acetic acid, 1%	2.61	2.63	2.61	2.43	2.17	0.87	0.14	0.01	< 0.01		
Hydro- chloric acid, 0.1%	0.11	0.02	< 0.01								
Phosphoric acid, 0.1%	< 0.01		—	—	—	—			—	—	
Trifluoro- acetic acid	1.20	0.78	0.54	0.34	0.22	0.06	< 0.02	< 0.01	—	—	
Diammo- nium phosphate, 50 mM	1.85	0.67	0.15	0.02	<0.01		_			—	
Triethy- lamine, 1%	2.33	2.42	2.50	2.45	2.37	1.96	0.50	0.12	0.04	< 0.01	
Buffers and	Salts										
Ammonium acetate, 10 mM	1.88	0.94	0.53	0.29	0.15	0.02	< 0.01				
Ammonium bicarbonate, 10 mM	0.41	0.10	0.01	< 0.01							
EDTA, disodium, 1 mM	0.11	0.07	0.06	0.04	0.03	0.03	0.02	0.02	0.02	0.02	
HEPES, 10 mM, pH 7.6	2.45	2.50	2.37	2.08	1.50	0.29	0.03	< 0.01		—	
MES, 10 mM, pH 6.0	2.42	2.38	1.89	0.90	0.45	0.06	< 0.01			—	

	Absor	Absorbance at specified wavelength (nm)									
	200	205	210	215	220	230	240	250	260	280	
Potassium phosphate, monobasic (KH ₂ PO ₄), 10 mM	0.03	<0.01									
Potassium phosphate, dibasic, (K ₂ HPO ₄), 10 mM	0.53	0.16	0.05	0.01	<0.01						
Sodium acetate, 10 mM	1.85	0.96	0.52	0.30	0.15	0.03	< 0.01			—	
Sodium chloride, 1 M	2.00	1.67	0.40	0.10	< 0.01						
Sodium citrate, 10 mM	2.48	2.84	2.31	2.02	1.49	0.54	0.12	0.03	0.02	0.01	
Sodium formate, 10 mM	1.00	0.73	0.53	0.33	0.20	0.03	< 0.01			—	
Sodium phosphate, 100 mM, pH 6.8	1.99	0.75	0.19	0.06	0.02	0.01	0.01	0.01	0.01	< 0.01	
Tris HCl, 20 mM, pH 7.0	1.40	0.77	0.28	0.10	0.04	< 0.01			—		
Tris HCl, 20 mM, pH 8.0	1.80	1.90	1.11	0.43	0.13	< 0.01	—				

Mobile phase absorbance measured against air or water (Continued)

	Absor	Absorbance at specified wavelength (nm)									
	200	205	210	215	220	230	240	250	260	280	
Waters [®] PIC [®] reagents											
PIC A, 1 vial/L	0.67	0.29	0.13	0.05	0.03	0.02	0.02	0.02	0.02	< 0.01	
PIC B6, 1 vial/L	2.46	2.50	2.42	2.25	1.83	0.63	0.07	< 0.01	—	—	
PIC B6, low UV, 1 vial/L	0.01	< 0.01								_	
PIC D4, 1 vial/L	0.03	0.03	0.03	0.03	0.02	0.02	0.02	0.02	0.02	0.01	
Detergents											
BRIJ 35, 1%	0.06	0.03	0.02	0.02	0.02	0.01	< 0.01		_	—	
CHAPS, 0.1%	2.40	2.32	1.48	0.80	0.40	0.08	0.04	0.02	0.02	0.01	
SDS, 0.1%	0.02	0.01	< 0.01	—	—	—	—	—	—	—	
Triton [®] X-100, 0.1%	2.48	2.50	2.43	2.42	2.37	2.37	0.50	0.25	0.67	1.42	
Tween [™] 20, 0.1%	0.21	0.14	0.11	0.10	0.09	0.06	0.05	0.04	0.04	0.03	

Mobile phase absorbance measured against air or water (Continued)

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