Gemini XPS
Dual-Scanning
Microplate Spectrofluorometer
Operator’s Manual

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Molecular Devices Corporation
Gemini XPS Operator’s Manual

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**Conventions Used in this Manual**

The names of keys that appear on the Gemini XPS control panel are shown in boxed Helvetica type. Example: `[Setup]`.

Italic and boldface type are used for emphasis. Examples: “Press *carefully* to engage,” “**Do not press down.**”

**NOTE:** A note provides information that will help you properly execute an action or procedure.

⚠️ **CAUTION:** Indicates an action or condition that could potentially damage the instrument or one of its components or could result in loss of data.

⚠️ **WARNING:** Indicates a situation that could result in potential injury to a person working with the system.

🔗 **BIOHAZARD:** Indicates a condition involving potentially infectious biological agents requiring that proper handling precautions be taken.
Chapter 1: Instrument Description

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Introduction

The Gemini XPS Dual-Scanning Microplate Spectrofluorometer can perform a variety of fluorescent applications as well as some time-resolved fluorescent and luminescent assays. The extreme flexibility and high sensitivity of the Gemini XPS make it appropriate for applications within the fields of biochemistry, cell biology, immunology, molecular biology, and microbiology.

The Gemini XPS uses two holographic diffraction grating monochromators which allow for individual optimization of wavelengths for both excitation and emission. The dual-scanning capability can also be used to determine excitation and emission settings for new fluorescent probes. Mirrored optics focus the light into the sample volume and cutoff filters are used to reduce stray light and minimize background interference. The light source is a high-powered Xenon flashlamp; additional flexibility is provided by allowing a variable number of lamp flashes per read.

The Gemini XPS also has two secondary modes that can be used for limited development of glow luminescence or time-resolved fluorescence assays. The performance of these two modes is not comparable to dedicated luminescence or time-resolved fluorescence instruments.

Microplates having 6, 12, 24, 48, 96, and 384 wells can be used in the Gemini XPS. Fluorescent detection is achieved from the top, through the entire depth of sample in the microplate well. Detection of species collected on membrane plates is also possible.

One plate carrier adapter is provided with the instrument. The adapter is required for optimum performance with standard 96- and 384-well format microplates for all applications.

The dynamic range of detection is from $10^{-6}$ to $10^{-11}$ molar fluorescein. Variations in measured fluorescence values are virtually eliminated by internal compensation for detector sensitivity, photomultiplier tube voltage and sensitivity, as well as excitation intensity.

Temperature in the microplate chamber is isothermal, both at ambient and when the incubator is turned on. When the incubator is on, the temperature may be controlled from 4°C above ambient to 45°C.

The contents of the wells in a microplate can be mixed automatically by shaking before each read cycle, which makes it possible to perform kinetic analysis of solid-phase, enzyme-mediated reactions such as a kinetic ELISA.

The Gemini XPS is controlled by an external computer running SoftMax® Pro software which provides integrated instrument control, data display, and statistical data analysis. The Gemini XPS cannot be operated without the computer and SoftMax Pro software.
Component Description

The main components of the Gemini XPS described in this manual are:

- The control panel
- The microplate drawer
- Optical system
- The back panel (connections and power switch)

Figure 1.1: The Gemini XPS

The Control Panel

Figure 1.2: Control Panel

The control panel consists of an LCD and four pressure-sensitive membrane keys which can be used to initiate and regulate the temperature and to open/close the drawer. When you press a control panel key, the Gemini XPS performs the desired action.

LCD

A 2×20-character liquid crystal display which shows the current instrument settings.
**Temperature Display**

The LCD panel displays the set point temperature and the actual temperature of the air inside the reading chamber.

**NOTE:** Allow 1 hour for the air temperature inside the reading chamber to equilibrate to the set point temperature to maintain the temperature of a microplate transferred from an incubator to the reading chamber.

**Keys**

Allows you to enter a set point at which to regulate the microplate chamber temperature. Pressing this key scrolls the temperature up or down, starting at the previous temperature setting (or the default of 37.0°C, if no setting had been made). Pressing the up (▲) or down (▼) arrow once increments or decrements the temperature shown in the display by 0.1°C; pressing and holding either arrow increments or decrements the temperature shown in the display by 1°C until it is released. If you increment the setting to the highest limit (45°C) and continue to press the up (▲) arrow, the display will not change. If you decrement the setting to the lowest limit, 15°C, and continue to press the down (▼) arrow, the display will not change.

Enables/disables the incubator function. When the incubator is on, the set temperature and actual temperature are shown on the front panel LCD display.

**NOTE:** When the instrument is performing a kinetic or spectral scan, the temperature keys on the front panel are disabled.

Opens or closes the microplate drawer. Whether or not the drawer will remain open depends on the incubator setting. If the incubator is off, the drawer will remain open; if the incubator is on, the drawer will close after approximately 10 seconds to assist in maintaining temperature control within the microplate chamber.

**NOTE:** To add reagents during a kinetic read, it is necessary to open the drawer by pressing the [Drawer] key. The drawer will only open, however, if the interval between readings is equal to the minimum read interval originally shown by SoftMax Pro software plus an additional 45 seconds. If you plan to open the drawer during a kinetic read, first determine the minimum read interval allowed and then increase the setting by a minimum of 45 seconds. The drawer will close automatically after this interval before the next read.
The Microplate Drawer

The microplate drawer, located on the right side of the Gemini XPS, slides in and out of the microplate chamber. A small plastic pusher, located in the front left corner of the drawer, holds the plate securely in place when the drawer is closed. The drawer remains in the reading chamber during read cycles.

One plate carrier adapter is provided with the instrument. The adapter is required for optimum performance with standard 96- and 384-well format microplates.

**NOTE:** The adapter must be removed to read 6-, 12-, 24-, or 48-well plates.

Microplate drawer operation varies, depending upon the incubator status. When the incubator is off, the microplate drawer remains open at power up and after a read. When the incubator is on, the drawer closes automatically to assist in controlling the temperature of the microplate chamber. To open the drawer, press the [Drawer] key. The drawer will remain open for approximately ten seconds, after which a beeping sound will alert you approximately two seconds before the drawer closes automatically. (The drawer will remain open longer during intervals set for adding reagents during kinetic reads—see the NOTE on page 1-5).

**NOTE:** Do not obstruct the movement of the drawer. If you must retrieve a plate after an error condition or power outage and the drawer will not open, it is possible to open it manually (see Chapter 5, “Troubleshooting”).

**Microplates**

The Gemini XPS can accommodate standard 6-, 12-, 24-, 48-, 96-, and 384-well microplates. Black-walled, clear-bottom or all black microplates are generally recommended for fluorescence assays because they have lower backgrounds than clear plates. White plates may be preferred for luminescence assays to optimize light collection.

Not all manufacturers’ microplates are the same with regard to design, materials, or configuration. Some plastics, most notably polystyrene, also have significant native fluorescence and can cause moderate to severe background fluorescence, especially in the UV range. If high sensitivity is required, it may be appropriate to use microplates that are designed to reduce background fluorescence.
The Optical System

1) The excitation light source is a xenon flashlamp. (Note that the lamp is off when luminescence mode is selected.)

2) The light passes through a band-pass filter which reduces the amount of stray light to the excitation monochromator.

3) The holographic diffraction grating monochromator selects the desired excitation wavelength.

4) The excitation beam is collimated by a mirror to a 1.0-mm diameter fiber before entering the sample in the microplate well. This focusing helps to prevent part of the beam from striking adjacent wells.

Figure 1.4: Components of the Gemini XPS Optical System
5) The light beam enters the well and, if fluorescent molecules are present, light of the emission wavelength is emitted back out to mirrors that focus it and send it to an optical bundle.

6) The emission monochromator (also a holographic diffraction grating monochromator) allows light of the chosen emission wavelength to pass to the emission filter wheel.

7) A long-pass filter further conditions the light prior to detection by the photomultiplier tube (PMT). This filter may be set automatically by the instrument or manually by the user.

8) The PMT detects the emitted light and passes a quantitative signal to the instrument’s electronics which then send the data to the computer.

The Back Panel

![Figure 1.5: Components on the Back Panel of the Gemini XPS](image)

The following components are located on the back panel of the Gemini XPS:

- **Power switch**: a rocker switch, labeled I/O (for on and off, respectively).
- **Power cord receptacle**: plug the power cord in here.
- **Fuse box cover**: cannot be opened while the power cord is plugged in. When opened, it provides access to the fuse box containing two fuses that are required for operation.
- **Parallel port**: Present but not used in this model.
- **Serial port** (double-shielded RS-232, for use with an external computer): plug one end of an 8-pin DIN serial cable into this port; the other end attaches to the serial (modem) port of the computer.
- **Labels**: provide information about the Gemini XPS, such as line voltage rating, cautionary information, serial number, etc. Record the serial number shown on this label for use when contacting Molecular Devices Technical Services.
Principles of Operation

Theory of Operation

Fluorescent materials absorb light energy of a characteristic wavelength (excitation) undergo an electronic state change, and instantaneously emit light of a longer wavelength (emission). Most common fluorescent materials have well-characterized excitation and emission spectra. Figure 1.6 shows an example of excitation and emission spectra for a fluorophore. The excitation and emission bands are each fairly broad, with half-bandwidths of approximately 40 nm, and the wavelength difference between the excitation and emission maxima (the Stokes shift) is typically fairly small, about 30 nm. Exceptions to this rule are the lanthanide dyes used in time-resolved fluorescence: they have a Stokes shift that is typically about 50 nm. There is considerable overlap between the excitation and emission spectra (gray area) when a small Stokes shift is present.

![Figure 1.6: Excitation and Emission Spectra](image)

Because the intensity of the excitation light is usually many tens of thousands of times greater than that of the emitted light, some type of spectral separation is necessary to reduce the interference of the excitation light with detection of the emitted light. The Gemini XPS incorporates many features designed to restrict interference from reflected excitation light. Among these features is a set of bandpass cutoff filters that may be set automatically by the instrument or manually by the user. If the Stokes shift is small, it may be advisable to choose an excitation wavelength that is as far away from the emission maximum as possible while still...
being capable of stimulating the fluorophore so that less of the excited light will overlap the emission spectrum, allowing better selection and quantitation of the emitted light.

Figure 1.7: Optimized Excitation and Emission Reading Wavelengths

Figure 1.7 shows that the best results are often obtained when the excitation and emission wavelengths used for reading are not the same as the wavelengths of the excitation and emission spectra of the fluorophore. When the reading wavelengths for excitation and emission are separated, a smaller amount of excitation light will pass through to the emission monochromator (gray area) and on to the PMT, resulting in a purer emission signal and more accurate data.

The Gemini XPS allows scanning of both excitation and emission wavelengths, using separate tunable monochromators. One benefit of being able to scan emission spectra is that you can assess more accurately whether the emission is, in fact, the expected fluorophore, or multiple fluorophores, and not one generated by a variety of background sources or by contaminants. Another benefit is that, if interfering fluorescent species are present, you may be able to find excitation and emission wavelengths that avoid interference.

For this reason, it may be desirable to scan emission for both an intermediate concentration of labeled sample, as well as the background of unlabeled sample. The optimum setting is where the ratio of the sample emission and background emission is at the maximum.

For more information regarding optimizing excitation and emission wavelengths using the spectral scanning capabilities of the Gemini XPS, please see “Optimizing Assays” on page 3-5.
Time-Resolved Fluorescence

In normal fluorescence mode, readings are taken while the lamp is on. The most common limitation to sensitivity in normal fluorescence is excitation energy or background fluorescence that cannot be eliminated from the emission signal. Since the lamp is the source of excitation energy, turning it off provides the best means of eliminating background excitation.

Time-resolved fluorescence is performed by flashing the excitation lamp and, after it is off, collecting the delayed emission for a period of time before the lamp is flashed again. Lanthanide dyes are frequently used to delay the fluorescence long enough to measure it after the lamp is turned off.

To assist with proper collection of data, you can also select when to start and end data collection (within the limits of the system—the minimum is 50 µsec and the maximum is 1450 µsec in 200-µsec steps).

Luminescence

In luminescence mode, no excitation is necessary as the species being measured emit light naturally. For this reason, the lamp does not flash, so no background interference occurs. A dark estimate is done over a dark reference and multiple readings are averaged together into one reading per well.

You can choose the wavelength where peak emission is expected to occur. In addition, multiple wavelength choices allow species with multiple components to be differentiated and measured easily. In luminescence read mode, no emission cut-off filter is used. The default setting for luminescence is the “zero order” position where the grating monochromator acts as a mirror which reflects all light to the PMT detector.

The Gemini XPS is a microplate spectrofluorometer with photomultiplier tube detection. Some luminescence applications like gene reporter assays may require a luminometer with photon counting detection for greater sensitivity.

Functional Description

The Gemini XPS is designed to be operated using SoftMax Pro software running on a computer connected to the instrument. Stand-alone functions are limited to setting and enabling temperature control and opening or closing the microplate drawer.

The information contained in this section provides an overview of the instrument capabilities. For a complete description of the modes of operation, how to choose instrument settings, etc., refer to the SoftMax Pro User’s Manual.
Modes of Operation

The Gemini XPS can perform three types of readings: fluorescence as well as some luminescence and time-resolved fluorescence applications. Within each read type, the Gemini XPS can perform four modes of operation: endpoint, kinetic, spectrum, and well scanning. Instrument setup parameters for each read mode are discussed in the SoftMax Pro User’s Manual.

**Endpoint Mode**

Select from one to four excitation/emission pairs in endpoint mode to obtain fluorescent (RFU), luminescent (RLU), or time-resolved fluorescent (RFU) readings for each well of a microplate. Endpoint data is reported as RFU/wavelength or RLU/wavelength.

**Kinetic Mode**

Kinetic analysis can be performed for up to 99 hours. The kinetic read interval depends upon the instrument setup parameters chosen in SoftMax Pro. At the end of a reading, rates are reported as RFU/second or RLU/second for each well.

Kinetic analysis has many advantages when determining the relative activity of an enzyme in different types of microplate assays, including ELISAs and the purification and characterization of enzymes and enzyme conjugates. Kinetic analysis is capable of providing improved dynamic range, precision, and sensitivity relative to endpoint analysis.

**Spectrum Mode**

Spectral analysis measures fluorescence or luminescence across a spectrum of wavelengths (minimum 250 nm; maximum 850 nm). When reading using fluorescence or time-resolved fluorescence, you may set a fixed wavelength for excitation and scan the emission wavelengths, and vice versa. The default value reported for each well is the wavelength of maximum fluorescence. When luminescence is chosen, only the emission wavelengths are scanned, and the default value reported for each well is the wavelength of maximum luminescence.

All spectrum readings are made using the scanning monochromators of the Gemini XPS.

**Well Scanning Mode**

Some applications that involve the detection of whole cells in large-area tissue culture plates may require the use of well scanning mode. As many cell lines tend to grow in clumps or in the corner of microplate wells, this non-confluent growth pattern may require multiple reads in a well at different locations.

When used with 6-, 12-, 24-, 48-, or 96-well plates, well scanning allows maximum surface area detection for whole cell assays. No plate adapter is required when using large-area tissue culture plates. For more information on well scanning, please review the appropriate section in the SoftMax Pro User’s Manual.
Temperature Regulation

The Gemini XPS has been designed to regulate the temperature of the microplate chamber from 4°C above ambient to 45°C. Upon power up, when the incubator is off, the temperature in the Gemini XPS microplate chamber is ambient and isothermal. Turning on the incubator by pressing the [Temp on/off] (incubator) key will cause the Gemini XPS to begin warming the microplate chamber. The temperature set point defaults to 37.0°C at start-up. With the incubator on, the temperature of the microplate chamber can be set and regulated from 4°C above ambient to 45°C.

NOTE: Accuracy of the temperature set point is only guaranteed if the set point is at least 4°C above ambient. If the temperature set point is lower than the ambient temperature, the chamber temperature will remain at ambient. Temperature regulation is controlled by heaters only and, therefore, cannot cool the temperature to a setting lower than ambient. Additionally, the highest setting (45°C) can be achieved only if the ambient temperature is >20°C.

You can change the temperature set point by pressing the up (\(^{\wedge}\)) or the down (\(\vee\)) arrow keys until the desired set point is shown above the key in the display.

The microplate chamber temperature is maintained at the set point until you press the incubator [Temp on/off] key again, turning temperature regulation off.

NOTE: Allow 1 hour for the air temperature inside the reading chamber to equilibrate to the set point temperature to maintain the temperature of a microplate transferred from an incubator to the reading chamber. If the incubator is turned back on after a momentary shut down, allow about ten minutes for the control algorithm to fully stabilize the microplate chamber reading.

Temperature regulation and control of the microplate chamber is achieved through electric heaters, a fan, efficient insulation, and temperature sensors. The heaters are located in the microplate chamber which is insulated to maintain the temperature set point. The sensors are mounted inside the chamber and measure the air temperature.

The temperature feedback closed-loop control algorithms measure the chamber air temperature, compares it to the temperature set point, and use the difference to calculate the regulation of the heating cycles. This technique results in accurate, precise control of the microplate chamber temperature with a temperature variation of the air across the entire microplate of less than 1°C. (Temperature uniformity within the microplate itself will depend upon its design, materials, and/or configuration.)
Automix

The Automix function permits automatic shaking of the microplate at preset intervals, thereby mixing of the contents within each well. Automix must be selected before beginning a reading. The actions associated with the Automix setting depends on the read mode chosen.

For endpoint mode, enabling Automix will shake the plate for a definable number of seconds and then read at all selected wavelengths.

When kinetic mode is chosen, two types of Automix can be enabled: you can set Automix to shake the plate for a definable number of seconds before the initial reading and/or for a definable number of seconds before each subsequent reading.

**NOTE:** Use of Automix is strongly recommended for ELISAs and other solid-phase, enzyme-mediated reactions to enhance accuracy.

Computer Control

The Gemini XPS is equipped with an 8-pin DIN RS-232 serial port through which the computer communicates with and controls the instrument (different types of cables are available for connecting to different types of computers—see Appendix A, “Cables and Accessories”). SoftMax Pro software, version 3.1 or greater, is required to control the Gemini XPS.
Specifications

**Fluorescence Photometric Performance**

- **Wavelength range:**
  - **Excitation** 250–850 nm
  - **Emission** 360–850 nm
- **Wavelength selection** Scanning monochromator tunable in 1-nm increments
- **Wavelength bandwidth** 9 nm
- **Wavelength accuracy** < ± 2.0 nm
- **Calibration** Self calibrating with built-in fluorescence calibrators
- **Sensitivity** (signal 3X STD DEV of baseline) 3.0 fmol/well FITC

**Time-Resolved Fluorescence Photometric Performance**

- **Data collection** 50–1450 µsec
- **Integration start/end** User selectable in 200-µsec intervals
- **Sensitivity** (signal 3X STD DEV of baseline) 0.5 fmol/well Eu-chelate (obtained with DELFIA reagent from Wallac Oy using a 384-well plate)

**Luminescence Photometric Performance**

- **Wavelength range** 360–850 nm
- **Sensitivity** (signal 3X STD DEV of baseline) 10 amol/well Alkaline Phos. (obtained with Emerald II reagent from Tropix, Inc.)

**General Photometric Performance**

- **Microplate formats** 6, 12, 24, 48, 96, 384
- **Light source** Xenon flash lamp (1 joule/flash)
- **Average lamp lifetime** 2 years normal operation (estimate)
- **Detector** Photomultiplier (R-3896)
- **Read time** 96 wells in 15 seconds; 384 wells in 45 seconds (measurement type may extend read time)
- **Dynamic range** 6 decades in 96-well black plates; auto PMT circuitry
- **Shaker** Time 0–999 seconds
- **Temperature control (chamber)** Ambient +4°C to 45°C

*NOTE:* Technical specifications are subject to change without notice.
General Photometric Performance (*continued*)

Sample evaporation control 90% RH compartment
Air temperature uniformity (across microplate) <1°C when temperature set point is 37°C
Ramp time to 37°C <1 hour

System Validation

Internal standards for fluorescence and wavelength

Environmental

Robot ready Yes
Turn-on time <5 min. to rated accuracy
Operating conditions 15 to 40°C
Operating humidity 0 to 90% RH non-condensing
Storage temperature -20 to 65°C

Physical

Size (h × w × d) 13.5" (340 mm) × 16.5" (420 mm) × 16.5" (420 mm)
Weight 35 lb (16 kg)
Power consumption 500 VA maximum
Line voltage and frequency 90–240 VAC, 50/60 Hz
Glossary of Terms

Automix
The Automix function determines how often, if at all, automated shaking of the microplate is performed during a reading. This feature is covered by U.S. Patent Number 5,112,134.

Cutoff
A filter used to condition the light entering or exciting the monochromators. In automatic mode, the instrument sets the cutoffs automatically based upon the wavelength(s) chosen for reading; with some read modes and types, however, you can choose a different filter wavelength (manual setting) for the emission monochromator.

Endpoint
A single reading made at one or more excitation/emission wavelengths.

Emission Spectral Scan
Measures fluorescence or luminescence across a spectrum of wavelengths for emitted light. The default value reported for each well is the wavelength of maximum fluorescence or luminescence.

Excitation Spectral Scan
Measures fluorescence at a single emission wavelength across a spectrum of excitation wavelengths. The default value reported for each well is the excitation wavelength of maximum fluorescence.

Fluorescence
The light emitted by certain substances resulting from the absorption of incident radiation. To measure fluorescence accurately, it is necessary to reduce light scatter. The governing equation for fluorescence is:

\[ \text{Fluorescence} = \text{extinction coefficient} \times \text{concentration} \times \text{quantum yield} \times \text{excitation intensity} \times \text{pathlength} \times \text{emission collection efficiency} \]

Fluorophore
A material which absorbs light energy of a characteristic wavelength, undergoes an electronic state change, and instantaneously emits light of a longer wavelength.

Gain
The amount of increase in signal power expressed as the ratio of output to input.

Incubator
Choosing Incubator from the Control menu or clicking the incubator button opens a dialog box allowing you to start or stop temperature regulation and to select an elevated temperature for the microplate chamber.

Instrument Setup
(In SoftMax Pro software) Defines the parameters (mode, wavelengths, automatic mixing, run time, read interval, etc.) used to read the microplate.
**Kinetic**
During kinetic readings, data is collected over time, with multiple readings made at regular intervals. The values calculated based on raw kinetic data are Vmax, Time to Vmax, and Onset Time. Kinetic readings can be single- or multiple-wavelength readings.

**LCD (Liquid Crystal Display)**
A 2×20-character display which shows the current instrument settings.

**Luminescence**
The emission of light by processes that derive energy from essentially nonthermal changes, the motion of subatomic particles, or the excitation of an atomic system by radiation.

**Parallel Port (Windows)**
A connection on a computer, usually LPT1, typically used with the cable for a parallel printer. Windows supports parallel ports LPT1 through LPT3.

**Photomultiplier Tube (PMT)**
A vacuum tube that detects light especially from dim sources through the use of photoemission and successive instances of secondary emission to produce enough electrons to generate a useful current.

**Read Mode**
The method used to read the microplate: endpoint, kinetic, spectrum, or well scan.

**Read Type**
The type of reading performed: fluorescence, time-resolved fluorescence, or luminescence.

**Readings per Well**
The number of times (settable by the user) that readings are taken on a well in fluorescence and time-resolved fluorescence modes. This setting also determines the amount of time that data is collected using the luminescence read type.

**SoftMax Pro**
An integrated software program (from Molecular Devices Corporation) that is used to control and collect data from the Gemini XPS instrument.

**Stokes Shift**
The difference between the wavelengths of the excitation and emission peaks.

**Time-Resolved Fluorescence**
Most fluorescence substances are not suitable for this type of reading. However, the fluorescence emitted by lanthanide dyes is delayed long enough to measure fluorescence after the lamp is turned off. Time-resolved fluorescence is used to reduce the amount of background noise which interferes with fluorescence. The excitation lamp flashes and, after it is off, the delayed emission is collected for a set period of time before the lamp is flashed again.
Chapter 2: Installation

Installation Warnings .......................... 2-3
Installation Cautions .......................... 2-3
Unpacking ....................................... 2-3
Setting Up the Instrument ..................... 2-3
Installing the Drawer Adapter ................. 2-4
Removing the Drawer Adapter ................ 2-5
**Installation Warnings**

1) Always make sure the power switch on the instrument is in the OFF position and remove the power cord from the back of the instrument prior to any installation or relocation of the instrument.

2) Do not operate the instrument in an environment where potentially damaging liquids or gases are present.

**Installation Cautions**

Do not touch or loosen any screws or parts other than those specifically designated in the instructions. Doing so might cause misalignment and could void the instrument warranty.

**Unpacking**

The Gemini XPS is packed in a specially designed carton. Please retain the carton and the packing materials. If the unit should need to be returned for repair, you must use the original packing materials and carton for shipping. If the carton has been damaged in transit, it is particularly important that you retain it for inspection by the carrier in case there has also been damage to the instrument.

**WARNING:** The Gemini XPS weighs approximately 35 pounds (16 kg) and should be lifted with care. It is recommended that two persons lift the instrument together, taking the proper precautions to avoid injury.

After examining the carton, place it on a flat surface in the upright position. Open the top of the box and lift the Gemini XPS, along with the packing materials around the ends, up and out of the shipping box. Remove the packing material from both ends of the instrument and set the instrument down carefully. The packing list that accompanies the instrument describes all components that should have been placed in the packing carton. Make sure all these items are present before proceeding.

**Setting Up the Instrument**

1) Place the Gemini XPS on a level surface, away from direct sunlight, dust, drafts, vibration, and moisture.

2) Turn the instrument around so that the back of the instrument is facing you as shown in Figure 2.1.
Figure 2.1: View of Rear Panel

3) Insert the female end of the power cord into the power receptacle at the rear of the Gemini XPS. Connect the male end to a grounded power outlet of the appropriate voltage. Molecular Devices recommends that you use a surge protector between the power cord and the grounded power outlet.

4) Insert the 8-pin DIN round end of the computer connection cord into the RS-232 serial port receptacle on the back panel of the instrument. Attach the other end to your computer (see Appendix A for more information).

5) Turn the Gemini XPS around so that the control panel now faces you. Be sure no cables run beneath the instrument. Leave at least three inches between the back of the instrument and the nearest objects or surfaces to ensure proper ventilation and cooling.

Installing the Drawer Adapter

If you are reading standard 96-well or 384-well microplates, you will need to install the drawer adapter.

⚠️ CAUTION: Incorrect insertion or removal of the adapter may cause damage to the microplate drawer of the Gemini XPS. The corner cut-out must be in the lower left corner where the plate pusher is located.

1) Turn power to the instrument on.

2) Press the [Drawer] button on the front panel or activate the drawer open command in SoftMax Pro software.

3) Hold the adapter so that the label is on the front side facing up.

4) Place the top back (Row A) portion of the adapter into the drawer first. The corner cut-out must be in the lower left corner where the plate pusher is located. While pushing against the back edge of the adapter, lower the front of the adapter into the drawer (see Figure 2.2).
Removing the Drawer Adapter

If the adapter is in the drawer and you are using “high profile” (6-, 12-, 24-, or 48-well) plates, you will need to remove the adapter.

⚠️ CAUTION: Incorrect insertion or removal of the adapter may cause damage to the microplate drawer of the Gemini XPS.

1) Turn power to the instrument on.
2) Press the [Drawer] button on the front panel or activate the drawer open command in SoftMax Pro software.
3) Remove the adapter plate.
Chapter 3: Operation

Prepare for a Reading. ............................ 3-3
  Turn the Instrument and Computer On .... 3-3
  Set the Temperature. ......................... 3-3
Read the Microplate. ............................ 3-3
Optimizing Assays .............................. 3-5
  Introduction. ................................. 3-5
  Using Spectral Scanning to Optimize Excitation and
  Emission Wavelengths for Fluorescence Assays. . . 3-6
Prepare for a Reading

Turn the Instrument and Computer On

The power switch for the Gemini XPS is located on the back panel. Press the rocker switch to the on position. The instrument will automatically perform diagnostic checks to ensure that it is functioning correctly. Turn the computer on at this time also and start the SoftMax Pro software program.

Set the Temperature

If elevated temperature within the microplate chamber is desired, you should turn on the incubator first, allowing enough time for the temperature to reach the set point before performing a reading. When you first turn the instrument on, up to 1 hour may be required for the temperature within the chamber to reach the set point. Turning on the incubator and choosing a temperature set point can be done using the software or the front panel of the instrument (described here).

**NOTE:** Temperature cannot be regulated at a set point that is lower than 4°C above the ambient temperature.

To enable the incubator, press the incubator [Temp on/off] key. The LCD display will indicate that temperature control is on and will show the set point and current temperature of the microplate chamber.

To change the temperature set point, press the up or down arrow keys until the desired temperature set point is shown in the display.

The microplate chamber temperature will be maintained at the set point until you disable temperature control by touching the incubator key again. When the incubator is off, the temperature within the microplate chamber will begin returning to ambient.

**NOTE:** Should you turn the incubator back on after a momentary shutdown, allow about ten minutes for the control algorithm to fully stabilize the microplate chamber temperature.

Read the Microplate

**BIOHAZARD:** The underside of the microplate must be dry prior to placing it in the drawer. If the microplate has fluid on the underside, dry it using a paper towel (or equivalent) before placing it in the drawer.

Insert the filled microplate into the drawer, matching well A1 with position A1 in the drawer. Make sure the microplate is flat against the drawer bottom.
(for 6-, 12-, 24-, or 48-well microplates) or against the adapter—see “Installing the Drawer Adapter” on page 2-4 for more information).

You must use SoftMax Pro software running on a computer (properly connected to the Gemini XPS) in order to perform instrument setup or reading and to analyze the data that is collected.

When reading is complete, the drawer of the instrument will open, allowing you to remove the microplate. If the incubator is on, the drawer will close again after approximately 10 seconds. If you return to the Gemini XPS and find the drawer closed after a reading has finished, press the [Drawer] key. When the drawer opens, you may remove the microplate.

**Operation Overview**

The following steps provide a quick reminder of the basic operating procedures required to perform an assay using the Gemini XPS.

1) Turn on the power switch of the Gemini XPS (located on the back panel). The microplate drawer will open automatically.

2) If you wish to regulate the temperature inside the microplate chamber, touch the [Temp on/off] (incubator) key to turn the incubator on and bring the microplate chamber to the default temperature of 37.0°C. The microplate drawer will close.

3) If the incubator is on, the LCD will show the current temperature of the reading chamber along with the temperature set point. To change the set point (to any setting from ambient +4° to 45°C), press the up or down arrow keys. Allow 1 hour for the air temperature inside the reading chamber to equilibrate to the set point temperature to maintain the temperature of a microplate transferred from an incubator to the reading chamber.

4) Select the desired instrument settings (read mode, type of analysis, template, etc.) using SoftMax Pro software on the external computer.

5) If you are performing kinetic analysis, add substrate at this time.

6) Load the prepared microplate into the drawer, being sure to match well A1 with the A1 mark on upper left-hand corner of the drawer. Use the drawer adapter, if required.

7) Use SoftMax Pro to start the reading.
Optimizing Assays

Introduction

The optimum instrument settings for detection of a particular fluorophore depend on a number of different factors. Settings that can be adjusted for assay optimization include the excitation and emission wavelengths, emission cutoff filter, readings per well, the PMT voltage, the temperature of the reading chamber, and the length of delay time for time-resolved fluorescence.

• The excitation and emission wavelengths may be set in 1-nm increments within the range of the instrument (250-850 nm). A procedure to optimize excitation and emission wavelengths for a given assay is outlined in the next section.

• The 14 emission cutoff filters assist in reducing background. Sources of background include stray excitation light and native fluorescence of plate materials, sample constituents, and solvents (including water). The default setting allows the instrument and SoftMax Pro software to determine which cutoff filter should be used (see Table 3.1 for default settings) in endpoint and kinetic modes. The spectral scan mode default uses no cutoff filter.

• The number of readings per well may vary between 1 (used for a quick estimate) and 30 (for very precise measurements). The default number of readings per well varies with the read mode: for fluorescence, the default is 6; for luminescence, the default is 30; for time-resolved fluorescence, the default is 20.

• The voltage of the photomultiplier tube may be set to low (for higher concentration samples), medium, or high (for lower concentration samples) in all read modes. In endpoint and spectrum mode, there is an additional setting, automatic, in which the instrument will automatically adjust the PMT voltage for varying concentrations of sample in the plate.

• The chamber of the Gemini XPS is isothermal at ambient as well as at elevated temperatures. The temperature in the reading chamber may be adjusted from 4°C above ambient to 45°C.

• In time-resolved fluorescence mode, you may set the integration start and end time in 200-µsecond increments (minimum 50 µsec/maximum 1450 µsec).

Other important factors that are independent of the instrument but which affect assay optimization include the Stokes shift. When the Stokes' shift is very small, optimizing the excitation and emission wavelengths and correct cutoff filter choices are very important.
Using Spectral Scanning to Optimize Excitation and Emission Wavelengths for Fluorescence Assays

1) Put 200 µL of sample that includes the fluorophore and 200 µL of a buffer control into separate wells of a microplate.

2) Excitation Scan
   A) Using SoftMax Pro, set up a plate section for a fluorescence read, spectrum mode, Em Fixed/Ex Scan, with no cutoff filter (default), and medium PMT.
   B) Set the emission wavelength based on the tentative value from the literature (or from a customary filter set used to measure your fluorophore). If the emission wavelength is not known, select a tentative emission wavelength about 50 nanometers greater than the absorbance maximum of the fluorophore. If necessary, the absorbance maximum can be determined by performing a spectral scan in a UV/Vis spectrophotometer.
   C) Set the excitation scan to start/stop approximately 50 nm below/above the tentative excitation value obtained from the literature (or the customary excitation filter).
   D) Set the step increment to 1 or 2 nm. (You may choose to do a preliminary scan with a 10-nm increment to determine the approximate peak location, and then repeat the scan over a narrower wavelength range with a 1 or 2 nm increment.)
   E) Perform the scan and view the results as a plot of emission fluorescence vs. excitation wavelength. Note the excitation wavelength at the emission peak and the maximum RFU value.

**NOTE:** If an error message reporting missing datapoints occurs, it may be due to possible saturation reported by SoftMax Pro at the end of the spectral scan. Reset the PMT to “low” and rescan the sample (scan the buffer blank with the PMT set to “medium” or “high”). If the error occurs after scanning with the PMT set to “low,” it may be necessary to dilute the sample.

**NOTE:** If the excitation scan shows no apparent peak, change the PMT setting to “high” and rescan the sample. If the spectral scan still shows no apparent peak, adjust the Y-scale of the zoom plot so that the plot fills the graph.

F) Select the optimal excitation wavelength. If the excitation peak wavelength and emission wavelength are separated by more than 80 nm, use the excitation peak wavelength value. If the excitation and emission wavelengths are less than 80 nm apart, use the shortest excitation wavelength that gives 90% maximal emission. (Follow the plot to the left of the peak until the RFU value falls to approximately 90% of the maximum and then drop a line from the 90% point on the plot to the x-axis—see Figure 3.1.)
3) Emission Scan #1
   A) In SoftMax Pro, set up a second plate section for a fluorescence read, spectrum mode, Ex Fixed/Em Scan, with no cutoff filter (default), and medium PMT.
   B) Set the excitation wavelength to the value determined in 2F above.
   C) Set the emission scan to start/stop approximately 50 nm below/above the tentative emission value obtained from the literature (or existing filter pair).
   D) Set the step increment to 1-2 nm (or do a preliminary scan with a 10-nm increment to determine the approximate peak location and then repeat the scan over a narrower wavelength range using a 1-2 nm increment.)
   E) Perform the scan and view the results as a plot of fluorescence vs. emission wavelength.

4) Select an emission cutoff filter that will block as much of the residual excitation light as possible without unduly reducing the fluorescence signal. The cutoff wavelength choices are 420, 435, 475, 495, 515, 530, 550, 570, 590, 610, 630, 665, or 695 nm. The cutoff value should be near the maximum emission wavelength (preferably between the excitation wavelength and the maximal emission wavelength) but at least 35 nm greater than the excitation wavelength.

5) Emission Scan #2
   A) In SoftMax Pro, set up a third plate section for an emission scan as specified in Step 3 above, except selecting Manual Cutoff Filter and setting the wavelength to that determined in Step 4.
   B) Perform the scan and view the results as a plot of fluorescence vs. emission wavelength. Note the wavelength giving the maximum emission (the optimal emission wavelength).
   C) Compare the spectra of the sample containing the fluorophore to the spectra of the buffer blank to get an estimate of the signal-to-noise ratio. If there is significant background interference, repeat steps 5A and 5B with another choice of cutoff filter.

---

**Figure 3.1: Plot of RFU vs. Wavelength**

- RFU at 90% of max λ
- max λ
- 90% of max λ
- Wavelength

<table>
<thead>
<tr>
<th>RFU at 90% of max</th>
<th>λ</th>
<th>max λ</th>
<th>λ 90% of max</th>
</tr>
</thead>
</table>
6) Results: The optimal excitation and emission wavelengths are those determined in steps 2F and 5B above.

7) Comments

A) In endpoint or kinetic fluorescence modes, the “Autofilter” feature will generally select the same cutoff filter wavelength as will the above optimization method. If desired, however, you may specify the cutoff filters manually.

B) For emission wavelengths less than 420 nanometers, experimental iteration is usually the best method of determining the optimal emission and excitation wavelengths. Begin optimization by performing steps 1–4 above. Try emission and excitation wavelength combinations with the 420 cutoff or with no cutoff filter. Similarly, for excitation wavelengths greater than 660 nanometers, try emission and excitation wavelength combinations with the 695 cutoff or with no cutoff filter.

Figure 3.2 shows the effects of different cutoff filters on a scan of fluorescein where excitation was fixed at 485 nm and emission was scanned from 490 nm to 560 nm (buffer blanks are not shown in this plot). Table 3.1 following lists default settings for the emission cutoff filters.

![Figure 3.2: Effects of Cutoff Filters on Fluorescein. Emission was scanned from 490 to 560 nm; excitation was fixed at 485 nm.](image-url)
### Table 3.1: Emission Cutoff Filter Default Settings

<table>
<thead>
<tr>
<th>Automatic Cutoff Selection</th>
<th>Endpoint and Kinetic Modes</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td>λ (nm)</td>
</tr>
<tr>
<td>1</td>
<td>None</td>
</tr>
<tr>
<td>2</td>
<td>420</td>
</tr>
<tr>
<td>3</td>
<td>435</td>
</tr>
<tr>
<td>4</td>
<td>455</td>
</tr>
<tr>
<td>5</td>
<td>475</td>
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<td>6</td>
<td>495</td>
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<td>8</td>
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<td>9</td>
<td>550</td>
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<td>10</td>
<td>570</td>
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<tr>
<td>11</td>
<td>590</td>
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<tr>
<td>12</td>
<td>610</td>
</tr>
<tr>
<td>13</td>
<td>630</td>
</tr>
<tr>
<td>14</td>
<td>665</td>
</tr>
<tr>
<td>15</td>
<td>695</td>
</tr>
</tbody>
</table>

Note: For spectrum mode, the default is “manual” (no automatic cutoff).
Chapter 4: Maintenance

Technical Support ........................................ 4-3
Cleaning ..................................................... 4-3
Cleaning the Fan Filter ................................. 4-4
Changing the Fuses ...................................... 4-4
Installing the Drawer Adapter ...................... 4-6
Removing the Drawer Adapter ...................... 4-7
Moving the Gemini XPS ............................... 4-8
Long-Term Shutdown .................................... 4-8
Technical Support

Molecular Devices Corporation is a leading worldwide manufacturer and distributor of analytical instrumentation. We are committed to the quality of our products and to fully supporting our customers with the highest possible level of technical service. In order to fully benefit from our technical services, please complete the registration card and return it to the address printed on the card.

If you have any problems using the Gemini XPS Dual-Scanning Microplate Spectrophotometer, in the U.S., contact our Technical Services group at 1-800-635-5577; elsewhere contact your local representative.

⚠️ WARNING: All maintenance procedures described in this manual can be safely performed by qualified personnel. Maintenance not covered in this manual should be performed only by a Molecular Devices representative.

⚠️ WARNING: Turn the power switch off and disconnect the power cord from the main power source before performing any maintenance procedure that requires removal of any panel, cover, or disassembly of any interior instrument component.

⚠️ WARNING: Removal of protective covers that are marked with the High Voltage warning symbol shown below can result in a safety hazard.

Cleaning

⚠️ BIOHAZARD: Wear gloves during any cleaning procedure that could involve contact with either hazardous or biohazardous materials or fluids.

Periodically, you should clean the outside surfaces of the Gemini XPS using a cloth or sponge that has been dampened with water. Do not use abrasive cleaners. If required, clean the surfaces using a mild soap solution diluted with water or a glass cleaner and then wipe with a damp cloth or sponge to remove any residue. Do not spray cleaner directly onto the instrument.

If needed, clean the microplate drawer using a cloth or sponge that has been dampened with water.

Should fluids spill in the drawer area (when the drawer is out), they will be directed to a tray at the bottom of the instrument, from which they will exit to the bench or counter beneath the instrument. Wipe up any spills immediately.

Never clean the inside of the instrument. Do not allow excess water or other fluids to drip inside the instrument.
Cleaning the Fan Filter

The fan filter on the bottom of the instrument requires periodic cleaning. The frequency of the cleaning depends on how dusty your particular lab is and could range from once a month to once every six months.

1) Turn power to the instrument OFF and then remove the power cord and cables from the back of the instrument.
2) Remove any plate or adapter from the instrument drawer. Turn the instrument over so that it rests flat on the bench.
3) Pop the black fan cover off and remove the filter.
4) Clean the filter by blowing clean, canned air through it or by rinsing it—first with water and then with alcohol—and allowing it to dry completely.
5) Place the clean, dry filter over the fan and replace the black cover.
6) Turn the instrument back over. Reconnect the power cord and cables to the instrument.

Changing the Fuses

Fuses burn out occasionally and must be replaced. If the instrument does not seem to be getting power after switching it on (the LCD shows no display), first check to see whether the power cord is securely plugged in to a functioning power outlet and to the receptacle at the rear of the Gemini XPS. If power failed while the Gemini XPS was already on, check that the power cord is not loose or disconnected and that power to the power outlet is functioning properly. If these checks fail to remedy the loss of power, follow the steps listed below to replace the fuses. Spare fuses (two U.S. and two metric) are shipped with the instrument. The U.S. and metric fuses are identical except for physical size. They may be taped to the back of the Gemini XPS.

If you no longer have spare fuses, you may obtain new ones from Molecular Devices (part numbers: 4601-0013 for U.S., 4601-0014 for metric) or from a local hardware store. Make sure fuses are rated SLOWBLOW (U.S.: 4-amp time-delay; metric: 4-amp, 5 × 20 mm, time-delay).

To change fuses, follow the steps below.

1) Switch power to the instrument off and then remove the power cord from the outlet and from the Gemini XPS power cord receptacle.
2) Remove the printer cable and computer cable (if connected) from the back of the Gemini XPS.
3) Turn the instrument around for easy access to the rear panel.
4) On the left-hand side of the rear panel (viewed from the back) is the power switch, fuse box, and power cord receptacle. As shown in the figures below,
press to the right of the black plastic cover of the fuse box to release it. Pull the fuse box cover away from the instrument. The fuse box will begin to slide forward.

![Fuse Box Cover](image)

**Figure 4.1: Power Switch, Fuse Box, and Power Receptacle**

5) Continue gently pulling the fuse box forward until it is free of the instrument.

![Removing the Fuse Box](image)

**Figure 4.2: Removing the Fuse Box**

6) When removed, the fuse assembly will appear as shown in Figure 4.3. The holder inside contains two fuses.

7) It is possible that only one of the fuses may have blown. Molecular Devices recommends that you replace both fuses, however, to ensure continued proper operation. Pull both fuses out of the holder and discard them.
Figure 4.3: The Fuse Box and Holder (with Fuses) Removed

8) Insert new SLOWBLOW-rated fuses into the fuse holder. Either end of the fuse may be forward.

9) Insert the fuse box into the opening in the instrument, making sure that the fuses are on the right side (toward the power receptacle). Press the fuse box into place, making sure the cover snaps closed.

10) Reconnect the power cord to the instrument and to the wall outlet and reconnect other cables previously disconnected.

Installing the Drawer Adapter

If you are reading standard 96-well or 384-well microplates, you will need to install the drawer adapter.

⚠️ CAUTION: Incorrect insertion or removal of the adapter may cause damage to the microplate drawer of the Gemini XPS. The corner cut-out must be in the lower left corner where the plate pusher is located.

1) Turn power to the instrument on.

2) Press the [Drawer] button on the front panel or activate the drawer open command in SoftMax Pro software.

3) Hold the adapter so that the label is on the front side facing up.

4) Place the top back (Row A) portion of the adapter into the drawer first. The corner cut-out must be in the lower left corner where the plate pusher is located. While pushing against the back edge of the adapter, lower the front of the adapter into the drawer (see Figure 4.4).
Removing the Drawer Adapter

If the adapter is in the drawer and you are using “high profile” (6-, 12-, 24-, or 48-well) plates, you will need to remove the adapter.

⚠️ CAUTION: Incorrect insertion or removal of the adapter may cause damage to the microplate drawer of the Gemini XPS.

1) Turn power to the instrument on.

2) Press the [Drawer] button on the front panel or activate the drawer open command in SoftMax Pro software.

3) Remove the adapter plate.
Moving the Gemini XPS

If you need to relocate the Gemini XPS, follow these steps.

⚠ **WARNING:** The Gemini XPS weighs approximately 35 pounds (16 kilograms). To avoid injury, it is recommended that two people lift the instrument together, using proper lifting techniques.

1) Remove any microplate (*and the adapter, if any*) from the drawer and then close the drawer.

⚠ **CAUTION:** Leaving the adapter in the drawer when moving the Gemini XPS could cause damage to the instrument.

2) Turn off the power switch and unplug the power cord from the source and from the receptacle on the back of the instrument.

3) Depending on the distance that you will be moving the instrument, you may wish to repackage the Gemini XPS in its original shipping carton. Otherwise, carry the instrument or place it on a rolling cart to transport it.

4) Ensure that the new location meets the proper specifications as described in Chapter 2, “Installation.”

Long-Term Shutdown

If you will not be using the Gemini XPS for an extended period of time, clean the *external* surfaces of the instrument.
Chapter 5: Troubleshooting

Error Codes and Resolutions ....................... 5-3
Opening the Drawer Manually .................. 5-4
This chapter lists error codes that may occur while using the instrument, followed by their most likely causes and remedies. Maintenance procedures are described in Chapter 4. For problems with the Gemini XPS that are not listed here, in the U.S., contact Molecular Devices Technical Services group at 1-800-635-5577; elsewhere, call your local representative.

**BIOHAZARD:** It is your responsibility to decontaminate the instrument, as well as any accessories, before requesting service by Molecular Devices representatives and before returning the instrument or any components to Molecular Devices Corporation.

**Error Messages**

The LCD will display *Fatal Error* codes when a situation arises that requires attention. Any reading in progress will stop. *Warning* messages do not stop a reading but are logged in the error buffer. Warning messages indicate a situation that requires attention but is not sufficient to stop or prevent a reading. Examples of situations that might cause warning messages are low memory, entries being out of range, or operations that could result in loss of data. These messages are generally self explanatory. For assistance regarding warning messages, contact your local Molecular Devices representative.

**Error Code Classifications**

Not all error messages are listed in this manual. The errors are grouped in relationship to possible causes as follows:

<table>
<thead>
<tr>
<th>Error Code Numbers</th>
<th>Possible Causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 - 199</td>
<td>Errors possibly caused by unrecognized commands being sent from the computer to the instrument.</td>
</tr>
<tr>
<td>200 - 299</td>
<td>Errors probably due to a main board failure or an error in the firmware code. Most of these errors require the assistance of Technical Support.</td>
</tr>
<tr>
<td>300 - 399</td>
<td>Instrument errors due to either a main board failure or other system failure. Most of these errors require the assistance of Technical Support.</td>
</tr>
<tr>
<td>400 - 499</td>
<td>Errors caused by a motor motion failure. Most of these errors require the assistance of Technical Support.</td>
</tr>
<tr>
<td>500 - 599</td>
<td>Errors due to failure or improper initialization of the instruments non-volatile memory (NVRAM). All of these errors require the assistance of Technical Support.</td>
</tr>
</tbody>
</table>

Some errors (shown in **boldface** in the following table) are considered fatal in that if they are detected during power up (the instrument will abort the power up sequence and display “FATAL ERROR ###” on the LCD panel). Check the following list to see if there is something that you can do to change the condition of the instrument to prevent the fatal error (e.g., closing the cuvette door during the power up sequence will prevent error numbers 111, 219, 302, and 310). Leave the instrument on for about five minutes, turn it off and then back on. If you continue to get the fatal error message on power up, record the error message number and contact Molecular Devices Technical Support or your local representative for assistance.
If the instrument is functioning normally when using SoftMax Pro, no errors should be in the buffer (except error number 100).

Table 5.1. Error Codes, Messages, and Notes about the Errors

<table>
<thead>
<tr>
<th>ERROR CODE</th>
<th>ERROR MESSAGE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>100-199: Operator Errors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>100</td>
<td>command not found</td>
<td>Command string not recognized.</td>
</tr>
<tr>
<td>101</td>
<td>invalid argument</td>
<td>Command Argument not recognized.</td>
</tr>
<tr>
<td>102</td>
<td>too many arguments</td>
<td>Too many arguments after command.</td>
</tr>
<tr>
<td>103</td>
<td>not enough arguments</td>
<td>Missing arguments.</td>
</tr>
<tr>
<td>104</td>
<td>input line too long</td>
<td>Too many characters in the input line.</td>
</tr>
<tr>
<td>105</td>
<td>command invalid, system busy</td>
<td>Instrument could not perform the give command because it was busy doing another task. Example: Request a wavelength while the monochromator is in motion.</td>
</tr>
<tr>
<td>106</td>
<td>command invalid, measurement in progress</td>
<td>Instrument could not perform command because a measurement was in progress</td>
</tr>
<tr>
<td>107</td>
<td>no data to transfer</td>
<td>Inputting transfer when there’s no data in the buffer</td>
</tr>
<tr>
<td>108</td>
<td>data buffer full</td>
<td>To many data sets in the buffer. Can be caused by setting up a long kinetic and disconnecting computer or SOFTmax PRO is preempted by another application.</td>
</tr>
<tr>
<td>109</td>
<td>error buffer overflow</td>
<td>More than 65 errors in the buffer, clear the buffer.</td>
</tr>
<tr>
<td>110</td>
<td>stray light cuvette, door open?</td>
<td>Cuvette door open while doing a read.</td>
</tr>
<tr>
<td>111</td>
<td>invalid read settings</td>
<td></td>
</tr>
<tr>
<td>200</td>
<td>assert failed</td>
<td>Firmware error.</td>
</tr>
<tr>
<td>200-299: Firmware Errors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>201</td>
<td>bad error number</td>
<td>Firmware error.</td>
</tr>
<tr>
<td>202</td>
<td>receive queue overflow</td>
<td>Caused by external device sending too much data over serial port and ignoring flow control.</td>
</tr>
<tr>
<td>203</td>
<td>serial port parity error</td>
<td>Parity bit error detected with incoming serial data.</td>
</tr>
<tr>
<td>204</td>
<td>serial port overrun error</td>
<td>Caused by host computer sending too much data and ignoring the flow control signal.</td>
</tr>
<tr>
<td>205</td>
<td>serial port framing error</td>
<td></td>
</tr>
<tr>
<td>206</td>
<td>cmd generated too much output</td>
<td>Firmware error.</td>
</tr>
<tr>
<td>207</td>
<td>fatal trap</td>
<td>Instrument error. Instrument locks up.</td>
</tr>
<tr>
<td>208</td>
<td>RTOS error</td>
<td>Firmware error.</td>
</tr>
<tr>
<td>209</td>
<td>stack overflow</td>
<td>Firmware error.</td>
</tr>
<tr>
<td>210</td>
<td>unknown interrupt</td>
<td>Firmware error.</td>
</tr>
<tr>
<td>300-399: Hardware Errors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>300</td>
<td>thermistor faulty</td>
<td>Unable to read a reasonable thermistor value. Thermistor faulty or disconnected, Main board problem, or ambient temperature out of range.</td>
</tr>
</tbody>
</table>
### Table 5.1. Error Codes, Messages, and Notes about the Errors

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<thead>
<tr>
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<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>301</td>
<td>safe temperature limit exceeded</td>
<td>A temperature of over 50°C detected on one or more of the 4 ther-</td>
</tr>
<tr>
<td></td>
<td></td>
<td>mistors. Temperature will be shut off and remain off until a suc-</td>
</tr>
<tr>
<td></td>
<td></td>
<td>cessful completion of power-up reset.</td>
</tr>
<tr>
<td>302</td>
<td>low light</td>
<td>Not enough light detected to make an accurate measurement. If doing</td>
</tr>
<tr>
<td></td>
<td></td>
<td>a cuvette read, the cuvette door may be open.</td>
</tr>
<tr>
<td>303</td>
<td>unable to cal dark current</td>
<td>Too much stray light detected on power-up, faulty or disconnected</td>
</tr>
<tr>
<td></td>
<td></td>
<td>pre-amp boards.</td>
</tr>
<tr>
<td>304</td>
<td>signal level saturation</td>
<td>During a cuvette read, could be due to cuvette door being open.</td>
</tr>
<tr>
<td>305</td>
<td>reference level saturation</td>
<td>During a cuvette read, could be due to cuvette door being open.</td>
</tr>
<tr>
<td>306</td>
<td>plate air cal fail, low light</td>
<td>Minimum signal/reference ratio not met during air calibration.</td>
</tr>
<tr>
<td>307</td>
<td>cuv air ref fail</td>
<td></td>
</tr>
<tr>
<td>308</td>
<td>stray light</td>
<td>Light leak in reading chamber or cuvette door open. Could also be</td>
</tr>
<tr>
<td></td>
<td></td>
<td>a faulty pre-amp board.</td>
</tr>
<tr>
<td>309</td>
<td>front panel not responding</td>
<td>LCD front panel bad or disconnected.</td>
</tr>
<tr>
<td>310</td>
<td></td>
<td></td>
</tr>
<tr>
<td>311</td>
<td></td>
<td></td>
</tr>
<tr>
<td>312</td>
<td>gain calibration failed</td>
<td>Power-up calibration and check of signal path gain is out of toler-</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ance. Could be due to bad or disconnected pre-amp or excessive</td>
</tr>
<tr>
<td></td>
<td></td>
<td>stray light.</td>
</tr>
<tr>
<td>313</td>
<td>reference gain check fail</td>
<td>Power-up check of the Reference amplifier’s gain out of tolerance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Could be due to bad or disconnected pre-amp board or excessive</td>
</tr>
<tr>
<td></td>
<td></td>
<td>stray light.</td>
</tr>
<tr>
<td>314</td>
<td>low lamp level warning</td>
<td></td>
</tr>
<tr>
<td>315</td>
<td>can’t find zero order</td>
<td>On power-up, grating motor could not find zero-order home</td>
</tr>
<tr>
<td></td>
<td></td>
<td>position.</td>
</tr>
<tr>
<td>316</td>
<td>grating motor driver faulty</td>
<td>Grating motor didn’t move to where it was commanded to in a</td>
</tr>
<tr>
<td></td>
<td></td>
<td>reasonable time.</td>
</tr>
<tr>
<td>317</td>
<td>monitor ADC faulty</td>
<td></td>
</tr>
</tbody>
</table>
Table 5.1. Error Codes, Messages, and Notes about the Errors

<table>
<thead>
<tr>
<th>ERROR CODE</th>
<th>ERROR MESSAGE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>400-499: Motion Errors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>400</td>
<td>carriage motion error</td>
<td>Carriage did not move to either of its photo interrupts in a reasonable time, or can't find its photo interrupt.</td>
</tr>
<tr>
<td>401</td>
<td>filter wheel error</td>
<td>Filter wheel did not move to its photo interrupt in a reasonable time, or can't find its photo interrupt.</td>
</tr>
<tr>
<td>402</td>
<td>grating error</td>
<td>Grating did not move to its photo interrupt in a reasonable time, or can't find its photo interrupt.</td>
</tr>
<tr>
<td>403</td>
<td>stage error</td>
<td>Stage did not move to its photo interrupt in a reasonable time, or can't find its photo interrupt.</td>
</tr>
<tr>
<td>500-599: NVRAM Errors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>500</td>
<td>NVRAM CRC corrupt</td>
<td>The CRC for the NVRAM data is corrupt.</td>
</tr>
<tr>
<td>501</td>
<td>NVRAM Grating cal data is bad</td>
<td>Grating calibration data is unreasonable.</td>
</tr>
<tr>
<td>502</td>
<td>NVRAM Cuvette air cal data error</td>
<td>Cuvette air calibration data is unreasonable.</td>
</tr>
<tr>
<td>503</td>
<td>NVRAM Plate air cal data error</td>
<td>Plate air calibration data is unreasonable.</td>
</tr>
<tr>
<td>504</td>
<td>NVRAM Carriage offset error</td>
<td>Carriage offset data is unreasonable.</td>
</tr>
<tr>
<td>505</td>
<td>NVRAM Stage offset error</td>
<td>Stage offset data is unreasonable.</td>
</tr>
<tr>
<td>506</td>
<td>NVRAM Battery</td>
<td>Time to replace the NVRAM battery (U3).</td>
</tr>
</tbody>
</table>

For all other error messages (codes not listed here), please contact your local Molecular Devices representative for assistance.

Opening the Drawer Manually

If an error occurs while the drawer is closed and you need to remove a microplate, press the [Drawer] key. If the drawer does not open, turn power to the instrument off and then on again. If the drawer remains closed, turn the incubator off (if it was on) by pressing the [Temp on/off] key.

If the drawer still remains closed, try using a blunt, flat object (such as a spatula) to open the door. With your index finger, pull the microplate drawer out of the instrument (do not force the drawer) and remove the microplate. This action will not harm the instrument, but should only be taken if the first two options have failed to open the drawer.

If you are still unable to open the drawer, contact your local Molecular Devices representative.
Appendix A  Cables and Accessories

Cables  . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . A-3
Items Available for Use with the Gemini XPS  . . . . . . A-3
Cables

Molecular Devices recommends that you use high-quality, double-shielded cables to connect the Gemini XPS to the computer. Choose cables that meet the following requirements:

Serial Interface Cable
(contact Molecular Devices for specific pin-out requirements)

Macintosh: Male DB8 to male DB8
IBM Compatible: Male DB8 to Female DB9

Items Available for Use with the Gemini XPS

Cable, RS-232 (instrument to Macintosh computer) ............. 9000-0091
Cable, RS-232 (instrument to PC computer) ............. 9000-0149
Power Cord ................................................................. 4400-0002
Fuse, 4-amp Time Delay ........................................... 4601-0013
Fuse, 4-amp (5 × 20 mm) Time Delay ......................... 4601-0014
Mouse Pad ............................................................... 9000-0133
Appendix B  Common Wavelengths
for Fluorescence, Time-Resolved
Fluorescence, and Luminescence

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Time-Resolved Fluorescence ..................... B-3
Luminescence. ........................................... B-3
**Common Wavelengths**

**Fluorescence**

<table>
<thead>
<tr>
<th>Fluorophore</th>
<th>Excitation Wavelength</th>
<th>Emission Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>HPPA</td>
<td>320</td>
<td>405</td>
</tr>
<tr>
<td>4-MeU, NADH, NADPH</td>
<td>355</td>
<td>460</td>
</tr>
<tr>
<td>Biotinidinase</td>
<td>355</td>
<td>544</td>
</tr>
<tr>
<td>PKU</td>
<td>390</td>
<td>485</td>
</tr>
<tr>
<td>Green Fluorescent Protein</td>
<td>390</td>
<td>510</td>
</tr>
<tr>
<td>Attophos/Attofluor</td>
<td>444</td>
<td>555</td>
</tr>
<tr>
<td>FITC</td>
<td>485</td>
<td>538</td>
</tr>
<tr>
<td>Ethidium Homodmer (DNA)</td>
<td>530</td>
<td>620</td>
</tr>
<tr>
<td>TRITC, Ethidium Bromide</td>
<td>544</td>
<td>590</td>
</tr>
<tr>
<td>Texas Red</td>
<td>584</td>
<td>612</td>
</tr>
<tr>
<td>TAMRA</td>
<td>547</td>
<td>580</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>280</td>
<td>340</td>
</tr>
<tr>
<td>La Jolla Blue</td>
<td>695</td>
<td>705</td>
</tr>
</tbody>
</table>

**Time-Resolved Fluorescence**

<table>
<thead>
<tr>
<th>Fluorophore</th>
<th>Excitation Wavelength</th>
<th>Emission Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eu-Chelate</td>
<td>360</td>
<td>610</td>
</tr>
</tbody>
</table>

**Luminescence**

<table>
<thead>
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<th>Probe</th>
<th>Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emerald and Emerald II&lt;sup&gt;a&lt;/sup&gt;</td>
<td>542</td>
</tr>
<tr>
<td>Sapphire and Sapphire II&lt;sup&gt;a&lt;/sup&gt;</td>
<td>461</td>
</tr>
<tr>
<td>Ruby&lt;sup&gt;a&lt;/sup&gt;</td>
<td>620</td>
</tr>
</tbody>
</table>

<sup>a</sup> Emerald, Emerald II, Sapphire, Sapphire II, and Ruby are trademarks of Tropix, Inc.
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