

# Typhoon 8600

## Instrument QuickStart v1.0

---

### Contents

#### Instrument guidelines

- Starting the instrument
- Shutting down the instrument
- Maintaining the instrument

#### Sample placement guidelines

- Dry samples
- Wet samples
- Sandwich gels

#### Scanner Control guidelines

- Setup recommendations
- Focal plane settings
- Press Sample option
- Orientation
- Multichannel fluorescence scans
- File structure
- Saving files

#### Scanning guidelines

- Previewing the scanned data
- Quickly assessing the quality of the scan

#### Getting information and Help

### Instrument guidelines

#### Starting the instrument

Turn on the Typhoon™ instrument and allow 30 minutes for the instrument to stabilize. The on/off switch is on the right side of the instrument. After the 30-minute warmup time, start the Scanner Control software and begin scanning.

#### Shutting down the instrument

Close the Scanner Control software. Then turn the switch on the right side of the Typhoon instrument to the off position.

#### Maintaining the instrument

To clean the glass platen and sample lid, dampen a lint-free cloth with distilled water and wipe the surface of the glass platen and sample lid. Alternatively, you can use a lint-free cloth dampened with 75% ethanol to wipe the surfaces, and then wipe the surfaces again using distilled water. Because laboratory alcohol formulations may contain residue that is highly fluorescent, make sure you follow alcohol cleaning with distilled water.

### Sample placement guidelines

#### Dry samples

Before you place a storage phosphor screen, microplate, dried membrane, or TLC plate on the glass platen, make sure the platen is clean and dry.

#### Wet samples

Place a small amount of distilled water on the glass platen. Then carefully position a wet gel (agarose or polyacrylamide) on the clean glass platen. Using too much water can cause the gel to move during the scan, which affects the quality of the collected data. In addition, do not trap air bubbles between the gel and the glass platen. Air bubbles will appear on the image.

#### Sandwich gels

You can collect data from polyacrylamide gels sandwiched between electrophoresis glass plates. To achieve optimum sensitivity, use low-fluorescence glass plates and make sure the bottom plate is 3 mm thick. To reduce diffraction patterns caused by the two different pieces of glass, use two Kapton™ strips (supplied in the Typhoon accessory kit) positioned over the spacers on the outside edges of the 3-mm thick plate to raise the sandwich gel slightly above the glass platen. Then use distilled water to fill the gap between the platen and the bottom of the 3-mm electrophoresis glass plate. If you use water, do not trap air bubbles between the sandwich gel and the glass platen. Rest one side of the sandwich gel on the glass platen and slowly lower it. When you can no longer lower the sandwich gel using your fingers, insert the Wonder Wedge™ tool (supplied in the Typhoon accessory kit) between the glass platen and the 3-mm electrophoresis glass plate. Then slowly remove the wedge. After scanning, use the Wonder Wedge to help remove the sandwich gel from the glass platen.

## Scanner Control guidelines

### Setup recommendations

Application	Focal Plane	Press Sample
Agarose Gel	Platen or +3 mm	No
Wet Polyacrylamide (PA) Gel	Platen	No
PA Sandwich Gel (3mm plate)	+3 mm	Optional
PA Sandwich Gel (<3mm plate)	Platen or +3 mm	Optional
Membrane or Blot	Platen	Optional
Microplate	+3 mm	Yes

### Focal plane settings

The Scanner Control software contains two settings for focal plane. The Platen setting focuses the optics to just above the glass platen. The +3 mm setting focuses the optics 3 mm higher than the glass platen. Choose the setting that best fits where the labeled target is positioned in the sample.

In general, use the Platen setting for thin samples (1 mm thick or less). Use the +3 mm setting for sandwich gels and microplates. For samples that can vary in thickness, such as agarose gels, scan test samples using both settings to determine which setting provides the best results.

### Press Sample option

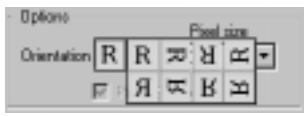
The Press Sample option prevents a sample from moving during scanning. Use the Press Sample option with minigel sandwich formats to hold the sandwich gel in place. Place a clean electrophoresis glass plate on top of a membrane or a blot sealed between plastic sheets or page protectors. (If necessary, you can use the Press Sample option to hold down the glass plate.) Do not use Press Sample with wet gels. Pressing a wet gel will damage the sample. Because a large polyacrylamide sandwich gel is heavy, you do not need the Press Sample option.

### Orientation

The A1 location for the sample placement is at the lower left corner of the glass platen. You can reduce the scan time by placing the long side of the sample along the lettered (A through R)

side of the glass platen. Use the orientation buttons to reorient the top of the sample. During the scan, an image of the sample appears in the ImageQuant™ Preview window.

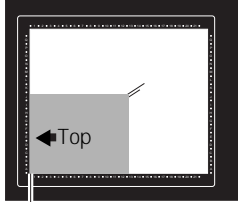
Orientation buttons





Screen or sample placement on the glass platen—



	For face-up sample, choose—	For face-down sample, choose—
--	-----------------------------	-------------------------------

---

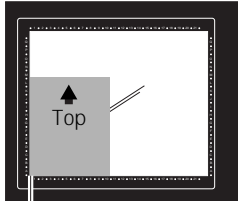


A1 corner











---

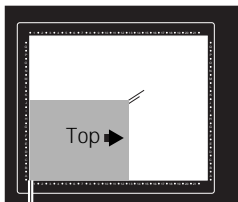


A1 corner











---

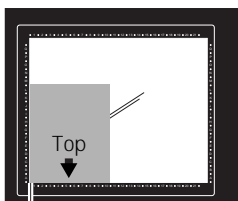


A1 corner






---



A1 corner

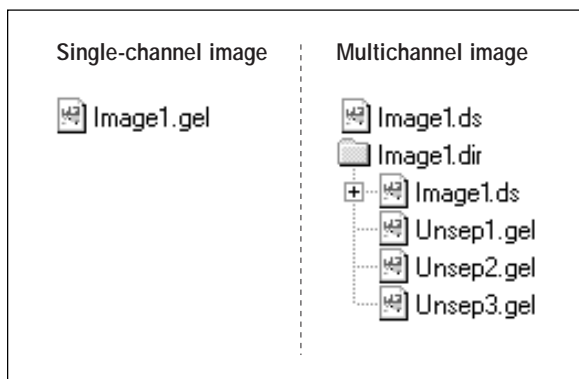
## Multichannel fluorescence scans

If all the following criteria are satisfied, two fluorescent scans can be acquired simultaneously:

- The same laser is selected.
- Different emission filters are selected.
- The same sensitivity setting is used.
- The beamsplitter wavelength is between the two emission filters and at least 5 nm from each emission filter.

## File structure

You can create two types of images using the Typhoon instrument and the Scanner Control software. All storage phosphor scans, most chemiluminescent scans, and fluorescent scans containing one dye create a single-image file, which is designated by the .gel file extension. Fluorescent scans that contain two, three, or four fluorescent dyes create a dataset image file, which is designated by the .ds file extension. A folder labeled with the .dir extension contains the individual .gel files that make up the dataset image. The folder also includes a backup copy of the .ds file.



## Saving files

To avoid the possibility of losing data, always save the image files on the local hard drive. After scanning, you can transfer the files and folders to a remote workstation and continue the analysis.

## Scanning guidelines

### Previewing the scanned data

The ImageQuant Preview window displays an image of the sample as the sample is scanned. You should monitor the preview image and check for saturated data. Saturated data appear as red areas in the image. If key areas of the image are saturated and you want to perform quantitation on the image, you will need to scan a fluorescent or chemiluminescent sample again using a lower PMT voltage setting. If you scanned a storage phosphor screen, you will need to expose a clean screen to the sample and reduce the length of the exposure time.

If the preview image displays all the relevant data before the instrument finishes the scan, you can cancel the scan and save the data. Doing this can eliminate scan time and reduce the image file size.

### Quickly assessing the quality of the scan

Display the scanned image in ImageQuant and use the Gray/Color Adjust, Pixel Locator, or Create Graph features to assess the signal values across the image.

## Getting information and help

- The *Typhoon User's Guide* provides step-by-step procedures for using the instrument and software.
- The Typhoon and Scanner Control online Help provides procedures for using the software.
- MD Scanner Technical Support provides assistance by phone or fax.

### **United Kingdom**

Amersham Pharmacia Biotech UK Limited  
Amersham Place Little Chalfont Buckinghamshire England HP7 9NA  
Telephone (44) 1494-544000, Fax (44) 1494-542266

### **United States and Canada**

Molecular Dynamics Inc  
Sunnyvale California USA  
Telephone (1) (800) 743-7782 or (1) (408) 773-1222, Fax (1) (408) 773-0152

### **Asia**

Amersham Pharmacia Biotech Asia Pacific Limited  
Taikoo Place Quarry Bay Hong Kong  
Telephone (852) 2811-8693, Fax (852) 2811-5251

---

ImageQuant, Molecular Dynamics, Typhoon, and Wonder Wedge are trademarks of Amersham Pharmacia Biotech Limited or its subsidiaries.

Amersham is a trademark of Nycomed Amersham plc.

Pharmacia and DropDesign are trademarks of Pharmacia and Upjohn Inc.

Kapton is a trademark of DuPont Corporation.

The Typhoon instrument is covered by one or more of the following U.S. patents: 5,528,050; 5,578,818; and foreign equivalents.

The Typhoon system is for research purposes only. It is not intended or approved for diagnosis of disease in humans or animals.

All goods and services are sold subject to the terms and conditions of sale of the company within the Amersham Pharmacia Biotech group that supplies them. A copy of these terms and conditions is available on request.

**Amersham Pharmacia Biotech UK Limited** Amersham Place Little Chalfont Buckinghamshire England HP7 9NA

**Amersham Pharmacia Biotech AB** SE-751 84 Uppsala Sweden

**Amersham Pharmacia Biotech Inc** 800 Centennial Avenue PO Box 1327 Piscataway NJ 08855 USA

**Amersham Pharmacia Biotech Europe GmbH** Munzinger Strasse 9 D-79111 Freiburg Germany

**Molecular Dynamics Inc** 928 East Arques Avenue Sunnyvale CA 94086 USA