iQ[™]5 Optical System Software – Instruction Manual

Compatible with the iQ5 and MyiQ™ Real Time PCR Detection System



NOTICE TO PURCHASER

This real-time thermal cycler is licensed under U.S. Patent No. 6,814,934 and corresponding claims in any counterpart Canadian patent thereof owned by Applera Corporation, solely in research and all other applied fields except human or veterinary in vitro diagnostics. No rights are conveyed expressly, by implication or estoppel to any patents on real-time methods, including but not limited to 5' nuclease assays, or to any patent claiming a reagent or kit.

Safety Information

Grounding

Always connect the iQ5 optics module to a three-prong, grounded AC outlet using the AC power cord provided with the system. Do not use an adapter to a two-terminal outlet. Always ensure that you set the module power switch to the off position when you connect or disconnect power cords.

Handling

Handle all components of the iQ5 system with care and with clean, dry hands at all times. The optical system contains mirrors and lenses that may shatter if the unit is dropped or struck with great force. If the unit is damaged such that internal components or wires are exposed, contact your local Bio-Rad office immediately. Do not attempt to repair or power on the instrument.

Servicing

The only user-serviceable parts of the iQ5 optics module are the lamp and filters. Call your local Bio-Rad office for all other optics module and thermal cycler related service. When you replace the lamp or filters, open only the outer casing of the iQ5 optics module. The camera lamp may get extremely hot during system operation. Do not attempt to remove the lamp without powering off the instrument and allowing the system to cool for at least 15 minutes. To prevent skin burns and fire hazards, do not attempt to operate the iQ5 system while the camera case is open. Do not open the casing of the iQ5 optics module when the instrument is in use.

Temperature

For normal operation, the maximum ambient temperature should not exceed 40°C. To ensure adequate cooling of the system, maintain a clearance of at least 4 inches around the sides of the iQ5 optics module. Do not block the fan vents near the lamp, as this may lead to improper operation or cause physical damage to the iQ5 detector. Do not operate the iQ5 optics module in extreme humidity (that is, greater than 90 percent) or where condensation can short internal electrical circuits or fog optical elements.

Notice

This Bio-Rad instrument is designed and certified to meet EN-61010 safety standards.

EN-61010 certified products are safe to use when operated in accordance with the instruction manual. This instrument should not be modified in any way. Alteration of this instrument will:

- Void the manufacturer's warranty
- Void the EN-61010 safety certification
- Create a potential safety hazard

Bio-Rad is not responsible for any injury or damage caused by the use of this instrument for purposes other than those for which it is intended, or by modifications to the instrument not performed by Bio-Rad or an authorized agent.

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Section 1 Introduction to the iQ[™]5 and MyiQ[™] Real-Time PCR Detection Systems

Nucleic acid amplification and detection are among the most valuable techniques used in biological research today. Real-time detection of PCR products is made possible by including in the reaction a fluorescent chemistry that reports an increase in the amount of DNA with a proportional increase in fluorescent signal. The fluorescent chemistries employed for this purpose include DNA-binding dyes and fluorescently labeled sequence-specific primers or probes.

Real-time analysis of PCR enables truly quantitative analysis of starting template copy number with accuracy and high sensitivity over a wide dynamic range. Specialized thermal cyclers equipped with fluorescence detection modules are used to monitor the fluorescence as amplification occurs. The iQ5 and MyiQ Real Time PCR Detection Systems build on the strengths of the iCycler[®] Thermal Cycler, which provides the optimum performance for PCR.

The iQ5 system features a broad-spectrum light source (tungsten-halogen lamp) that offers maximum flexibility in selecting fluorescent chemistries. The filter-based optical design of the iQ5 allows the use of optimal wavelengths of light for excitation and emission, resulting in excellent sensitivity and discrimination between multiple fluorophores. The MyiQ optical design supports the use of a single filter pair optimized for excitation and emission of green fluorescent dyes, resulting in excellent sensitivity for the detection of fluorophores such as FAM and SYBR Green I. In both systems, a CCD detector captures a simultaneous image of all 96 wells. This results in a comprehensive data set illustrating the kinetic behavior of the data during each cycle.

Simultaneous image collection insures that well-to-well data may reliably be compared. The iQ5 software reports data on the PCR in progress in Real Time, allowing immediate feedback on reaction success. All of these features demonstrate that the hardware of the iQ5 & MyiQ systems was built to promote reliability and flexibility.

The iQ5 software includes features that make software easy and useful. The software is designed for convenience-offering speedy setup and analytical results. The functions are presented graphically to minimize hunting through menus. Tips on usage are available as your mouse glides over the buttons. In addition, the Online Help Manual is available at all times simply by pressing the F1 key. The iQ5 software automatically analyzes the collected data at the touch of a button yet leaves room for additional optimization of results based on your analysis preferences.

1.1 Setting Up the System Hardware

The iQ5 or MyiQ system should be installed on a clean, dry, and level surface. Identify an appropriate work area for the installation process prior to unpacking any system components. The entire installation process should take approximately 15 minutes to complete. Handle the optics module and iCycler thermal cycler with care and with clean, dry hands during unpacking and assembly. Do not handle any system components with wet hands.

1.1.1 System Checklist

- Optics Module (170-9751: iQ5, or 170-9744: MyiQ), 1 box
- Power cord
- iQ5 software installation disk (170-9753)
- Amplification tech notes CD (version 1.0)
- iCycler Chassis (170-8701), 1 box
- Power cord
- iCycler chassis
- iCycler thermal cycler operating instructions
- iQ5 Optical Kit (170-9752), 1 box
- Optical reaction block
- Modified sliding rear cover for iCycler thermal cycler
- Serial cable
- USB cable
- Filter extraction tool
- Optical tape applicator tools (3)

- iQ5 optics support bracket
- Support bracket screws
- Hex driver
- Hex screws
- Spare part: halogen lamp
- iQ Calibration Kit (170-8792), shipped on dry ice
- External well factor solution
- 1x calibration dye solutions (9 dyes, 3 tubes of each dye)

Contact your local Bio-Rad office if any system components are missing or damaged.

Accessories

In addition, the following accessories are required to complete the installation process:

- Scissors
- Optical-quality sealing film or tube caps
- #2 Phillips screwdriver
- Calibrated micropipet(2)
- Optical-quality PCR plates or tubes
- Aerosol barrier pipet tips

1.1.2 Installing the Optics Module on the iCycler Chassis

- 1. Remove the existing rear cover from the iCycler chassis by sliding the cover towards the front of the iCycler base.
- 2. Install the modified sliding rear cover provided with the optical reaction block, ensuring that the notch is oriented towards the rear of the iCycler thermal cycler.
- 3. Push the sliding rear cover on top of the chassis as far back as possible.
- 4. Rotate the green latches on the optical reaction block up towards the open lid.
- 5. Lift the optical reaction block by the handle and install it onto the chassis. Lower the front portion of the reaction block so that it engages with the chassis before the rear portion. The rear of the block lid should fit over the front lip of the sliding rear cover.
- 6. Secure the optical reaction block in place by rotating the green latches downward.
- 7. Close the optical lid.

1.1.3 Installing the Support Bracket

A support bracket with roller is provided for the iQ5 or MyiQ optics module. It is mounted to the rear of the iCycler thermal cycler.

Align the optics module support bracket with the two holes on the rear of the iCycler thermal cycler.

Using a #2 Phillips screwdriver, adjust the height of the bracket with two of the appropriate screws provided with the system accessories. Both of the screws should be approximately in the center of the slots on the bracket.

1.1.4 Installing the Optics Module

- 1. Remove the plastic sheath and protective label from the optics module and place the optical module on a flat surface, taking care not to touch the inside of the nose portion of the module.
- 2. Remove the protective label from the optical lid.
- 3. Slide the optics module onto the U-bracket.
- 4. Secure the optics module to the U-bracket using the two long, thin hex screws and the hex driver provided.
- 5. After the optics module has been installed, confirm that the optics module and lid assembly can be opened and closed readily.

If the lid is difficult to open, lower the support bracket slightly before tightening the bracket mounting screws.

If the lid is difficult to close, try raising the support bracket slightly before tightening the rear screws.

1.1.5 Connecting Power and Communication Cables to the System

Before connecting any communication or power cables to the system, confirm that the power switch for the iCycler thermal cycler and the optics module are in the OFF position.

1. Close the optical reaction block by sliding the lid forward and pressing down on the lid handle. On the right side of the optics module are three connectors.

2. Using the cables provided in the iQ5 or MyiQ Optical Kit, establish power and communication with the computer as follows:

- Recessed 3-pin power connector Connect the supplied power cord between the optics module and a grounded power outlet. This connection provides power only to the optical module; a separate power cord must be connected to the iCycler thermal cycler.
- Serial connector A serial connector is located at the rear of the iCycler thermal cycler. Connect the serial cable to the rear of the iCycler chassis and to the serial port on the side of the optics module. This connection enables communication between the iQ5 software and the iCycler thermal cycler.
- USB port connector Connect the supplied USB cable between the optics module and a USB 2.0 high-speed enabled port on the computer. Data are transferred to the computer via this cable. This single connection directs the operation of both the optics module and the iCycler thermal cycler by the iQ5 Optical System Software. At the right rear corner of the optical reaction block is a single connector:
- Positive docking connector This self-aligning connector is secured into place when the optics module is installed on the iCycler chassis. This connection senses when the lid handle is lifted.

1.2 Installing the iQ5 Software

Locate the software installation disk provided with the iQ5 system. This installation disk is compatible with computers running the Windows XP and/or Windows 2000 operating systems.

- 1. Insert the iQ5 Optical System Software installation CD in a CD-ROM drive.
- 2. If the installation program does not begin automatically, click Run... in the Start menu and then type X:\iQ5\Setup, where X is the drive letter of the CD-ROM drive. For example, if the CD-ROM is the E drive, type E:\iQ5\Setup.
- 3. Follow all screen prompts to finalize the installation. Certain configurations of Windows 2000 and XP initialize new folders by assigning Read and Execute permission for the members of the Users group. If you have this type of operating system and this is a first-time installation, the administrator must change the protection for the Program Files/Bio-Rad folder or the Program Files/Bio-Rad/iQ5 folder so that you can save protocol, plate setup, and data files.
- 4. If, after you change the protection on either of these folders, you still cannot write to the folders beneath the Program Files/Bio-Rad/iQ5 folder, check the properties of each folder. Specifically, in the Properties window Securities tab, ensure that the check box that allows inheritable permissions to propagate to that folder has been selected.

Confirm that the iQ5 system software is working properly by double-clicking on the shortcut icon located on the Windows desktop, or by selecting the iQ5 program icon from the Bio-Rad folder in the Windows Start menu.

1.2.1 Camera Driver Installation

Before using the real time PCR detection system for the first time, five procedures must be performed on the new instrument: Camera Driver Installation, Mask Alignment, Background Calibration, Persistent Well Factor Collection, and Pure Dye Calibration.

Note: Pure Dye Calibration is **NOT** performed on the MyiQ real time PCR detection system.

- 1. Power on the iCycler thermal cycler and the optics module.
- 2. Windows will display a "Found New Hardware Wizard" dialog box. To install the iQ5 drivers, select the option to "Install from a list or specific location". Click Next to continue.
- 3. In the new window that appears, select the option to "Search for the best driver in these locations", and select the "Include this location in the search" checkbox. Use the browse button to navigate to the "iQ5\Drivers" folder. The default location of the drivers is: C:\Program Files\Bio-Rad\iQ5\Drivers. Click Next to continue.

- 4. Next Windows will display a Hardware Installation dialog box regarding Windows Logo Testing, a service offered by Microsoft. To complete driver installation for the iQ5 or MyiQ systems, click on the "Continue Anyway" button.
- 5. Allow the instrument optics to warm up for 20 min prior to performing the Mask Alignment, Background Calibration, Persistent Well Factor Collection, or Pure Dye Calibration procedures. Please refer to Section 2.1 for additional information on system calibration.

1.2.2 Overview of the iQ5 Software

The iQ5 software is divided into five sections, called "modules". Icons representing each of the modules are always shown on the left side of the screen. The active or selected module has an orange background, whereas unselected modules have a gray background. Each module is subdivided into windows that perform a specific function for that module.

The five modules in the iQ5 software are the:



Workshop Module: This module is used to select a Plate and Protocol and Run an experiment. It is also where experimental files are selected and opened for Data Analysis. The Workshop Module consists of a Setup and Plate Summary window. The Protocol, Plate, Run Set and Data File tabs which can be used to select, open, edit or create files. In Setup the selected Protocol and Plate Setup can be Run or the selected data file can be opened for analysis in Data Analysis.



Run-Time Central Module: This module is used to initiate and monitor experimental runs. It is accessed from the Workshop Module, once the Protocol and Plate Setup have been chosen, by clicking Run or Run End Point.



Data Analysis Module: This module contains a suite of tools enabling you to conduct thorough and varied analyses of your experimental data. Within this module are screens for Quantitative, Melt Curve/Peak, End Point, Allelic Discrimination, and Gene Expression analyses. The Edit Plate screen permits post-run editing of the experimental plate setup, allowing you to correct erroneous sample type assignments. The Data Analysis module is opened automatically when you open a saved data file from the Workshop.



Calibration Module: In order to extract the best data from your real time PCR experiment, the iQ5 must be calibrated. These simple and easily-performed calibration routines are accessed in the Calibration module. There are calibration routines for pure dyes, performed when you add a new fluorophore to your suite, Mask Alignment, background and well factors.

User Profile

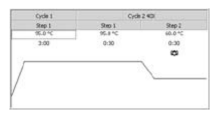
User Profile Module: This module can be used, by an iQ5 Administrator, to add new Users and to set Users access/restrictions to various functions of the iQ5 software.

All Users can use this module to set their personal preferences for the iQ5 software.

Section 2 Quick Guides to Running Experiments on the iQ5 or MyiQ System

To set up a real-time PCR experiment:

1. Select, Edit or Create a Protocol.



2. Select, Edit or Create a Plate Setup.

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3. Click Run.

Run

4. Run the Selected Protocol and Plate Setup.



2.1 Calibration Quick Guide

Every experiment on the iQ5 and MyiQ systems requires that the instruments are calibrated. You calibrate the iQ5 and MyiQ cameras using the iQ5 software Calibration Module.

There are four calibrations required for the iQ5 device which must be done in the following order:

- 1. Mask Alignment
- 2. Background Calibration
- 3. Persistent Well Factor Generation
- 4. Pure Dye Calibration

There are three calibrations required for the MyiQ device, which must be done in the following order:

- 1. Mask Alignment
- 2. Background Calibration
- 3. Persistent Well Factor Generation

NOTE: Pure Dye Calibration is NOT performed on the MyiQ real time PCR detection system

This order is the order that the Calibration tabs occur in the iQ5 Software (from left to right). Choose the **Mask Alignment** tab first and then when you complete each calibration choose the tab to the right of the completed calibration to do the next calibration.

Calibration factors are specific for the optical path you are using which includes the Reaction vessel, sealing mechanism and volume used in the experiment you wish to run.

2.1.1 Calibration Overview

You need the following to calibrate the iQ5 camera:

- 1. iCycler iQ Calibrator Dye Solution Set
- 2. iCycler iQ External Well Factor Solution (found in the iQ Calibrator Dye Solution Set)
- 3. 3 x 96-well PCR plate or preferred reaction vessel
- 4. Optical quality sealing tape or preferred sealing method

With these materials, you must prepare three plates: the external well factor plate, the background calibration plate, and the pure dye calibration plate.

Preparation of an External Well Factor Plate

(For Mask and Persistent Well Factor calibrations)

To prepare an external well factor plate:

- 1. Dilute the 10x External Well Factor Solution to 1x with water.
- 2. In Plate 1 pipet equal volumes of the 1x external well factor solution into ALL wells.
- 3. Seal the plate using the sealing method you will use in your experiment.

Preparation of an Background Calibration Plate

Prepare a background calibration plate by sealing Plate 2 using the sealing method you will use in your experiment.

Preparation of the Pure Dye Calibration Plate

To prepare a pure dye calibration plate:

- 1. In Plate 3, pipet equal volumes of the appropriate 1x calibrator solution into 8 wells of the 96-well plate, using the volumes that you will use in your experiment.
- 2. Repeat for all fluorophores needed. See example Pure Dye Plate Setup
- 3. Seal the plate using the sealing method you will use in your experiment.

2.1.2 Mask Alignment

Before you align the mask, complete the following steps:

- 1. See Calibration Overview about preparing the external well factor plate.
- 2. Place the external well factor plate in the iCycler instrument.
- 3. Select the Mask tab in the Calibration module.

To align the mask:

- 1. Click Home.
- 2. Click the Filter Position 2 option button.
- 3. Click Take an Exposure.
- 4. Click Show Mask.
- 5. If any pink pixels are present in the mask, reduce the exposure time and retake an exposure. Keep retaking the exposure until no pink pixels are present.
- 6. Click Optimize Mask.
- 7. Click Save Mask.

This completes Mask Alignment calibration. Next, perform Background Calibration.

2.1.3 Background Generation

Before you perform background calibration, complete the following steps:

- 1. See Calibration Overview about preparing the background calibration plate.
- 2. Ensure that the mask has been aligned.
- 3. Place the Background calibration plate in the iCycler instrument.
- 4. Select the Background tab in the Calibration module.

To perform background calibration:

- 1. Select a well seal.
- 2. Select a vessel type.
- 3. Click Collect Background Factors.
- 4. When iQ5 completes the Background Calibration run, a dialog box appears with the message, "Background Calibration Run Complete."
- 5. Click OK to exit.

This completes background calibration. Next, perform Persistent Well Factor calibration.

2.1.4 Persistent Well Factor Calibration

Before you perform a Persistent Well Factor Calibration run, complete the following steps:

- 1. See Calibration Overview about preparing the external well factor plate.
- 2. Ensure that the mask has been aligned and that Background Calibration has been performed.
- 3. Place the external well factor plate in the iCycler instrument.
- 4. Select the Well Factors tab in the Calibration module.

To perform a Persistent Well Factor Calibration run:

- 1. Select a sample volume.
- 2. Select a well seal.
- 3. Select a vessel type.
- 4. Click Collect Persistent Well Factors.
- 5. When iQ5 completes the Persistent Well Factor Calibration run, a dialog box appears with the message, "Persistent Well Factor Calibration Run Complete".
- 6. Click OK to exit.

This completes Persistent Well Factor calibration. This completes calibration of MyiQ cameras. For iQ5 cameras, perform Pure Dye Calibration as described below.

2.1.5 Pure Dye Calibration

Before you perform a Pure Dye Calibration run, complete the following steps:

- 1. See Calibration Overview about preparing a Pure Dye Calibration plate.
- 2. Ensure that the mask has been aligned.
- 3. Ensure that Background Calibration has been performed
- 4. Ensure that Persistent Well Factors have been generated.
- 5. Place the Pure Dye Calibration Plate in the iCycler instrument.
- 6. Select the Pure Dye tab in the Calibration module.

To perform a Pure Dye Calibration run:

- 1. Select or Create a Pure Dye Plate Setup.
- 2. Click Run Pure Dye Calibration.
- 3. When iQ5 completes the Pure Dye Calibration run, a dialog box appears with the message, "Pure Dye Calibration Run Complete".
- 4. Click OK to exit.

This completes calibration of the iQ5 Camera.

Note: SYBR and SYBR1-5 do not require calibration.

2.1.6 Viewing Calibration Files

View	Reports	Tools
Ap	plication Lo)gs
Dal	ta File Log	
Dal	ta File Info	1
Cal	libration Da	ata
An	alysis Para	meters

Calibration files can be viewed by selecting Calibration Data from the View menu.

Background factors file will be found in the Background Factors folder in the iQ5 Program Folder.

Persistent Well Factors will be found in the Well Factors folder in the iQ5 Program Folder.

Pure Dye Calibrations will be found in the RMEData folder in the iQ5 Program Folder

2.2 Protocol Quick Guide

Protocols can be:

- 1. Selected,
- 2. Created, and
- 3. Edited from the Protocol Viewer of the Workshop module.

2.2.1 Selecting a Protocol

To select a protocol.

- 1. Locate the folder by using the browser below the Protocol button.
- 2. Click the protocol name. The selected protocol appears in the Selected Protocol window.

Note: The iQ5 software has a number of sample protocol files which may be used.

2.2.2 Editing or Creating a Protocol

- Edit a selected protocol by clicking Edit in the Selected Protocol window.
- Create a protocol from a protocol template by clicking Create New in the Selected Protocol window.

Note: Clicking Edit or Create New in the Selected Protocol window opens the Protocol Editor.

Edit the protocol change the appropriate fields in the spreadsheet by performing one or more of the following five tasks:

1. Edit the Dwell Time and Setpoint temperature.

Click in the Dwell Time or Setpoint cell, and then enter the Dwell Time or Setpoint temperature. To enter 10 seconds, type 0 followed by: then 10 (that is, as the time appears in the spreadsheet). Alternatively 10 seconds can be entered as 0.10.

2. Acquire data.

Click in the Data Acquisition column, and then click Real-Time at the step you want to collect real-time data. Click Melt Curve if data from a melt curve is required.

Note: Ensure that every thermal protocol has at least one Data Acquisition step.

3. Insert cycles and steps.

Insert a cycle by clicking in the Insert column within the cycle row. Cycles have a blue background. the iQ5 software inserts the new cycle below the current cycle.

Insert a step by clicking in the Insert column within a step row. Steps have a white background. the iQ5 software inserts the new step below the current cycle.

Note: You can use the Options cell to customize whether the step or cycle is inserted before or after the current cycle/step, as well as how many steps the iQ5 software will insert when you insert a cycle.

4. Delete cycles and steps.

Delete a cycle by clicking in the Delete column within a cycle row. Cycles are indicated with a blue background.

Delete a step by clicking in the Delete column within a step row. Steps are indicated with a white background.

5. Save the protocol.

Click Save & Exit Protocol Editing. Enter the name of the protocol in the Save As dialog box, and then click Save.

Note: You can only exit the Protocol Editor by clicking Save & Exit Protocol Editing or Cancel & Exit Protocol Editing.

2.3 Plate Setup Quick Guide

To set up a plate:

- 1. From the Workshop module:
 - a. Click Create New in the plate setup display window to enter the Plate Setup Editor.

or

b. Click Plate. Open the desired plate setup file in the appropriate directory. Double click the file name to go directly to the Plate Setup Editor.

or

c. Click Plate. Open the desired plate file in the appropriate directory. Click the file name to open the plate setup in the bottom right pane of the Workshop window. Click Edit to open the plate setup in the Plate Setup Editor.

or

- d. Click Data File. Open the desired data file in the appropriate directory. Click the file name to open the plate setup associated with the data file in the bottom right pane of the Workshop window. Click Edit to open the plate setup in the Plate Setup Editor.
- 2. Enter or edit any notes about the plate setup in the Notes box.
- 3. Enter or edit the sample volume, seal type, and vessel type.
- 4. Enter or edit a name for the experiment.
- 5. Click Select/Add Fluorophores and select or edit the fluorophores to be used on the plate. Associate each fluorophore with a unique color.
- 6. Clear the Whole Plate Loading check box if it is not applicable; otherwise, leave the check box selected. If you are editing a plate, the Whole Plate Loading check box may be unavailable because it is not appropriate based on the current definition of the plate.

- 7. Click a sample type icon.
- 8. Select the type of replicate loading desired.
- 9. Click a fluorophore.
- 10. Click or drag across the plate to define wells with the selected fluorophore and sample type.
- 11. Continue defining the remaining wells that will contain the first fluorophore by changing to the other sample type icons.
- 12. To delete a previously defined well, click the Delete All icon, and then click the well.
- 13. To delete the selected fluorophore from a previously defined well, click the Delete Fluorophore icon, and then click the well.
- 14. If there are standards in the first dye layer, click Dilution Series and calculate the concentrations of the standards, or type the concentrations in the spreadsheet manually

2.4 Running a Real-Time Experiment on the iQ5 or MyiQ System

After you click Run in the Workshop module, the iQ5 software opens the Initiate Run tab within the Run-Time Central module.

2.4.1 Beginning a Run

To begin a run:

- 1. Check that the desired Protocol and Plate Setup are displayed in the bottom half of the Initiate Run screen.
- 2. Select the type of well factors to use by selecting one of the following two check boxes:



Fig. 2.2. The Well Factor Dialog Box.

- 3. Click Begin Run.
- 4. Name the file in the Save Optical Data File dialog box.
- 5. Click OK.

2.4.2 During a Run

When the iQ5 software begins the run, it opens the Monitor Run window. You can see the progress of the run in this window.

At the end of the run, the Run Status dialog box appears as shown below. You can choose between displaying the data in the Data Analysis module or returning to the Workshop Module.

Run Status		X
Completed Display completed run in data analysis?	(Yes)	No

Fig. 2.3. The Run Status dialog box.

Click Yes to proceed to the Data Analysis module.

2.5 Data Analysis Quick Guide

The Data Analysis module is where data is presented and analyzed. When the iQ5 software opens, the Data Analysis module is gray and the module is not active. To analyze a data file, open the data file in the Data File tab of the Workshop module by selecting the data file and then clicking Analyze.

The Data Analysis module consists of six tabs:

PCR Quant

The PCR Quantification tab is used to set the analysis conditions for the data file. The analysis conditions include setting the PCR baseline, the threshold and which wells to exclude or include in the experiment. The analysis conditions should be set before using the Gene Expression, End Point or Allelic Discrimination tabs. For experiments with standards of known quantities, the PCR Quant tab is also where absolute quantities can be determined. The efficiency of the PCR reaction can also be determined using standard curves with either known quantities used to produce the standard curve or by using a serial dilution of the template under investigation.

• Melt Curve/Peak

Melt Curve/Peak is a dynamic tool used to measure the melting temperature (T_m) of double stranded DNA molecules. DNA duplexes can be visualized by either incorporation of DNA-binding dyes (for example, SYBR Green I) or by hybridization with fluorescently labeled probes. Three major applications for Melt Curve/Peak are peak identification (number of amplified products), characterization of molecular beacons, and allelic discrimination.

End Point

End Point Analysis provides a convenient method of analyzing final RFU (Relative Fluorescence Unit) values. This can be useful when PCR Analysis is to determine if a given sample is Positive or Negative for a particular nucleic acid sequence.

Allelic Disc

The Allelic Discrimination tab is useful for assigning genotypes to unknown samples by making comparisons to known genotypes. It can be used to distinguish among homozygous wild types, homozygous mutants and heterozygous individuals based either on threshold cycle or on RFU.

• Gene Expr

The Gene Expression screen has flexible tools for the determination of the fold induction of one gene relative to another gene or relative to itself under different circumstances, e.g., temporally, geographically or developmentally different points.

Edit Plate

In the Edit Plate screen, you may make changes to the sample type assignment or to the quantities of the Standards in the plate setup used to run the experiment. This is a simple way to salvage your experiment should you make mistakes in the sample type assignment. While you may make changes to the plate setup, the original plate setup is never discarded and always remain with the data file so that you may revert to it at any time.

2.5.1 PCR Quant Tab Quick Guide

The PCR Quant screen is where conditions are set for the analysis of amplification data. Once the wells to be used in the analysis are selected and the baseline and threshold set, the software determines the threshold cycle for each well and, if standards are present, estimates the starting quantity of each unknown sample. These calculations take place for each dye layer present in every well. The software will automatically set the baseline individually for each well and then automatically calculate a single threshold value for each dye layer. If standards are present, the software searches for a threshold that gives the statistically best standard curve. The baselines and threshold may also be set manually.

Use the PCR Quant tab to set the analysis conditions for the data file. The analysis conditions include:

- Setting the PCR baseline.
- Setting the threshold.
- Determining which wells to exclude or include in the experiment.

To analyze a data file:

- 1. Click the Data File tab in the Workshop module.
- 2. Select a data file, and then click Analyze. The file opens in the PCR Quant tab within the Data Analysis module. iQ5 automatically chooses the Data Analysis conditions including baselines and thresholds.
- 3. You may select or deselect wells to be included in the analysis by selecting the Analyze Wells check box.
- 4. If the data open in any screen other than the PCR Quant screen, click PCR Quant. If the data file is being opened for the first time, an automated analysis of baselines and threshold will be conducted, including every defined well. If the file was previously saved after an analysis, the last set of analysis conditions are applied again.
- 5. Make any manual adjustments to the baselines.
- 6. Make any manual adjustments to the threshold.
- 7. You may return to the automated analysis at any time by choosing Baseline Threshold from the context menu accessed by a right click on the amplification chart.

2.5.2 End Point Quick Guide

You can implement End Point Analysis in two ways:

- Click Run End Point to initiate the collection of End Point data from a sample plate.
- Click the Endpoint tab in the Data Analysis module for an existing data file.

To initiate an end point run:

- 1. Insert the experimental plate into the iCycler reaction module.
- 2. Select or Create the Plate Setup in the Workshop module Plate Setup tab.
- 3. Click Run End Point.
- 4. In the Run-Time Central/Initiate Run tab, specify the setpoint for data collection, and then click Begin Run.

Note: You must use Persistent Well Factors for every End Point Run.

- 5. Enter a name for the data file, and then click Save.
- 6. Once iQ5 completes the run, the End Point tab is displayed.
- 7. Make selections for the following parameters:
 - Method Use Negatives to differentiate samples that do not amplify the target sequence from those that do
 amplify the target sequence.
 - End Point Tolerance and Tolerance Parameter
- 8. Select the wells to analyze by clicking Analyze Wells.
- 9. Define the positive and/or negative controls in the Define Controls column within the End Point Analysis table.
- 10. Click Recalculate. The End Point analysis table displays a positive, negative, or blank label for each unknown under the Unknowns Call column.
- 11. Click Reports to obtain customized reports of the End Point analysis.

End Point Analysis of an Existing Data File

- 1. Select the data file to be analyzed in the Workshop module Data Analysis tab.
- 2. Click Analyze.
- 3. Click the End Point tab.
- 4. Make selections for the following parameters:
 - Method

- End Point Tolerance and Tolerance Parameter
- 5. Select the wells to analyze by checking the Analyze Wells button. Click on Analyze Selected Wells, and close the Select Wells for Analysis floating window when finished.
- 6. Define the positive and/or negative controls in the Define Controls column of the end point analysis table.
- 7. Click Recalculate. The End Point analysis table displays a positive, negative, or blank label for each unknown under the Unknowns Call column.
- 8. Click Reports to obtain customized reports of the End Point analysis.

2.6 Allelic Discrimination Quick Guide – For Multiplex Data Only

The Allelic Discrimination feature of the Data Analysis module is available post-run and offers flexibility for analyzing allelic discrimination data from multiplex PCR experiments. You can display samples on a scatter plot based on threshold cycle or RFU (relative fluorescence units) values at any PCR cycle. You can have the iQ5 software automatically make allele calls, or you may manually make the allele calls.

2.6.1 Analyze an Allelic Discrimination File

To analyze an allelic discrimination file:

- 1. Click on the Data File tab in the Workshop module.
- Select a data file, and then click Analyze. The file opens in the PCR Quant tab of the Data Analysis module. iQ5 automatically chooses the Data Analysis conditions. To manually adjust these conditions, refer to the PCR Quant Tab.
- 3. You can select or deselect wells that will be included in the analysis by selecting the Analyze Wells check box.
- 4. Click the Allelic Discr tab in the Data Analysis module.
- 5. Select the Fluorophores that represent Allele1 and Allele2 in the Assign Fluorophores area as shown below.

Assign Fluorophores X - Axis Allele (1) Fluorophore	TET	-
- Axis Allele (1) Hoor ophore	J.C.	
/ - Axis Allele (2) Fluorophore	TexasRed	-
	1	
	Switch Fluoro	phores

Fig. 2.4. The Assign Fluorophores Dialog Box.

6. Choose between the Threshold Cycle and RFU display modes in the Display Mode area, shown below, to display the allelic discrimination data.

Di	play Mode	
	Threshold Cycle	
(RFU	

Fig. 2.5. The Allelic Discrimination Display Mode Selection Box.

• Threshold Cycle displays the distribution of samples on the scatter plot based on the threshold cycle.

Note: Samples that do not cross threshold will be assigned the threshold cycle value of the last cycle run in the experiment.

• RFU displays the distribution of samples on the scatter plot based on the RFU generated by each sample at the last PCR cycle number. Click the drop-down list next to the Select Cycle box to generate the scatter plot based on an RFU from a different cycle of the PCR experiment.

7. Click Automatic Call or Manual Call, as shown below. Automatic Call is the default parameter.



Fig. 2.6. The Allelic Discrimination Analysis Mode Selection Box.

a. In Automatic Call, threshold bars are positioned automatically in one of two ways, as shown in the example below:

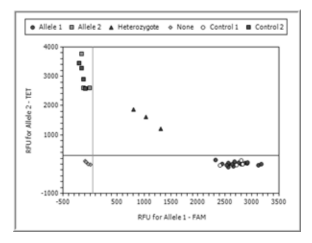


Fig. 2.7. The Allelic Discrimination Results Graph.

- Based on distribution of the control wells, when at least three wells have been assigned to Control 1 and three to Control 2.
- At 90 percent of the threshold cycle range or 10 percent of the RFU range on each axis if no controls are named.
- You may click and drag threshold bars directly on the plot to adjust Automatic Call assignment.
- b. Manual Call is the alternative analysis mode. Adjustments may be made either in the scatter plot or in the data spreadsheet.
 - Scatter plot: Make a selection in the Allele Call box. Then click and drag the cursor over the corresponding sample(s) in the scatter plot. iQ5 reassigns the samples to the allele call selected by the radio button, and updates the data spreadsheet accordingly.
 - Data spreadsheet: Click in the Call cell of the spreadsheet. A menu appears that lets you select an allele call for that sample. Once you select an allele call, the scatter plot reflects the change.
- 8. Click Reports in the menu to obtain customized reports for the allelic discrimination data.

2.7 Gene Expression Quick Guide

In order to make valid comparisons between two genes, their measured expression levels must be normalized in some way. Consider the example of an experiment in which one well is loaded with 100 ng of cDNA and another with 1 ng of cDNA. There is an immediate bias by a factor of 100 if the resultant data are taken at face value.

Normalization can be part of the qPCR experiment by inclusion of a reference gene, presumed to have constant levels of expression over the variation under study. Otherwise, normalization takes place before the qPCR using some other measurement such as cell number, RNA mass, or tissue mass.

The iQ5 software can present expression data normalized to one or more reference genes, or, for data normalized before PCR, as a relative quantity. The final interpretation of relative quantitation data may require additional calculation outside the iQ5 software. For example, when normalization is based on number of cells and sample A comes from twice as many cells as sample B, if both A and B have identical threshold cycles, the software will report a relative quantity of A/B = 1. To obtain the correct relationship, divide by the normalizer, cell number. Since A started with twice as many cells as B, the normalizer for A:B is 2. Applying the normalization factor changes the final result to show the A is expressed at half the level of B.

2.7.1 Relative Quantity (ΔC_T) Quick Guide

To calculate Relative Quantity (ΔC_T):

- 1. In the PCR Quant tab, establish desired experimental conditions such as baseline and threshold and which wells are to be included in the analysis.
- 2. Click the Gene Expression tab.
- 3. Select Relative Quantity as your analysis method.
- 4. Assign Gene Names and Condition Names in the Gene Expression Plate interface.
- 5. Assign attributes (sample color, show graph, etc.) in the Gene List.
- 6. Assign attributes (sample color, show graph, etc.) in the Condition List.
- 7. Click Recalculate to see your results.

If you need to compare these data to results obtain in other .opd files, you will need to enable this file for Multi-file Gene Expression analysis, also called a Gene Study. To enable your file for Gene Study:

- 8. Click the Enable for Gene Study button.
- 9. Go to the File menu to save your file.

Relative Quantity is graphed in the Gene Expression Graph Interface. You can access the data in the Gene Expression Settings Interface in the Data Table.

2.7.2 Normalized Expression ($\Delta\Delta C_T$) Quick Guide

To calculate Normalized Expression ($\Delta\Delta C_T$):

- 1. In the PCR Quant tab, establish desired experimental conditions such as baseline and threshold and which wells are to be included in the analysis.
- 2. Click the Gene Expression tab.
- 3. Select Normalized Expression as your analysis method.
- 4. Define Genes and Conditions in the Gene Expression Plate interface, if necessary.
- 5. Assign attributes (sample color, show graph, etc.) in the Gene List.
- 6. Assign attributes (sample color, show graph, etc.) in the Condition List.
- 7. Click Recalculate to see your results.

If you need to compare these data to results obtain in other .opd files, you will need to enable this file for Multi-file Gene Expression analysis, also called a Gene Study. To enable your file for Gene Study:

- 8. Click the Enable for Gene Study button.
- 9. Go to the File menu to save your file.

Normalized Expression is graphed in the Gene Expression Graph interface. The data can be accessed in the Gene Expression Settings interface within the Data Table.

2.7.3 Multiple File Gene Expression Quick Guide

To create a Gene Study (from multiple .opd files):

- 1. Set PCR Quant analysis conditions for each .opd file.
 - a. In the PCR Quant tab, establish desired experimental conditions such as baseline and threshold and which wells are to be included in the analysis.
 - b. Click the Gene Expr tab.
 - c. Make sure all files to be included in the Gene Study has the Enable for Gene Study button active.
 - d. Save each individual file.
- 2. From the menu toolbar select File/New/Gene Study.
- 3. In the Gene Expression Study Manager select Add OPDs.
- 4. Select the files that you wish to add to the Gene Study. More than one file can be selected.
- 5. Once the files are added to the Gene Expression Study Manger, select OK.
- 6. Select your analysis method (either Normalized Expression or Relative Quantity).
- 7. Assign Gene and Condition Names in the Gene Expression Plate Interface.
- 8. Assign Attributes in the Gene List.

Note: In Normalized Expression analysis you must assign reference gene(s).

9. Assign Attributes in the Condition List.

10. Click Recalculate to see your results.

Normalized Expression is graphed in the Gene Expression Graph Interface. The data can be accessed in the Gene Expression Settings Interface within the Data Table.

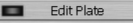
2.8 Edit Plate Setup Quick Guide

For post-run editing of the Plate Setup saved in a Data File:

- 1. From the Workshop module:
 - a. Click Data File above the directory of the home workshop. Navigate the directory until the desired data file is found. Double click the file name to bring the file directly into the Data Analysis module.

or

- b. Click Data File above the directory of the home workshop. Navigate the directory until the desired data file is found. Click the file name once to open the plate setup associated with the data file in the bottom right pane of the Workshop window. Click Analyze to bring the data to the Data Analysis module.
- 2. At the top of the Data Analysis window, click Edit Plate.



3. A modified version of the Plate Setup Editor will open. In this modified window, you cannot add fluorophores to or remove fluorophores from the fluorophore list. Nor may you add a previously-defined fluorophore or remove a previously-defined fluorophore from a well.

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ł	С		•		1	2	3	4	5	6			1	●Cy5	Cy5			
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Fig. 2.8. The Edit Plate Window.

- 4. Edit any notes about the plate setup in the Notes box.
- 5. Edit the name of the experiment in the Experiment Name box.
- 6. Deselect the Whole Plate loading option button if it is not applicable. Otherwise, leave this option button selected. If Whole Plate Loading is unavailable when you begin editing the plate, you may not restore Whole Plate Loading.

To edit a sample type.

- a. Click a Fluorophore.
- b. Click the correct sample type icon.
- c. Select the Next # check box to be sure that the desired number will be assigned to the next standard you define.



- d. Select the type of Replicate loading desired.
- e. Click a previously defined well to overwrite its original sample type definition. Replicates of the well are not affected.
- f. Repeat steps b-e above for each additional sample type to be edited for this fluorophore.
- g. Repeat steps a-f above for each additional fluorophore.

To edit the concentrations of the standards:

- a. Click the fluorophore for the standards to be defined.
- b. Click Dilution Series and calculate the concentration, or enter it directly in the spreadsheet.
- c. Select the Units for the standards.

Click Apply Plate Changes to make the changes. To see the effect on analysis, go to one of the other Data Analysis windows.



The original plate setup is retained with the data file and may be restored at any time by clicking Restore Original Plate.

Section 3 Workshop Module

The Workshop module consists of two tabs: the Setup tab and the Plate Summary tab. The primary tab in the Workshop module is the Setup window, as described below.

3.1 Assay Setup Window Layout

With the Setup tab active, the displayed window contains four additional tabs:

- **Protocol**: Used to select, edit, or create a protocol for running a real-time experiment.
- Plate: Used to select, edit, or create a plate setup for running a real-time experiment.
- **Run Set**: Consists of a linked protocol and plate setup. This tab is useful when you repeat the same experiment on a regular basis.
- **Data File**: Primarily used to select and open a data file. You can also use this tab to run a real-time experiment using the same protocol and plate setup that were used to create the data file. The selected plate setup can be either the Original plate setup or the Current (last saved) plate setup.

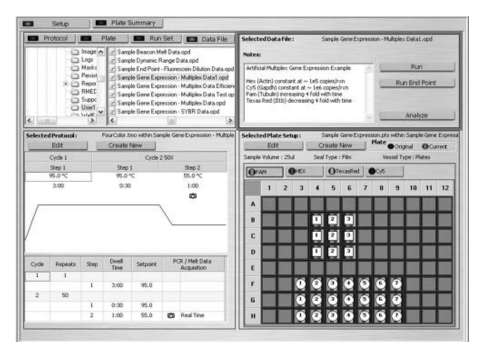


Fig. 3.1. The Setup tab.

The Workshop Setup window consists of four areas.

- 1. **File Browser area**: Located in the top left of the Setup window, the File Browser area contains a folder tree on the left side and a file list on the right side. Between the folder tree and file list is a moveable dividing bar. Moving this bar can provide more space in either the folder tree or the file list, which can help with navigation.
 - The files displayed in the file list depend upon the selected Setup window tab.
 - If the Protocol tab is selected, the iQ5 software displays protocol files, which have the extension .tmo. The selected protocol appears in the Selected Protocol window.
 - If the Plate tab is selected, the iQ5 software displays plate setup files, which have the extension .pts. The selected plate appears in the Selected Plate Setup window.

- If the Run Set tab is selected, the iQ5 software displays run set files, which have the extension .run. The iQ5 software displays the selected run set, which is a linked protocol and plate setup, in the Selected Protocol and Selected Plate Setup windows.
- If the Data File tab is selected, the iQ5 software displays data files created from previous experimental runs, which have the extension .opd. The iQ5 software displays the selected data file name and any associated notes in the Selected Data File window. The Notes box is editable only after you open the data file. The iQ5 software displays the protocol and plate setup used in creating this data file in the Selected Protocol and Selected Plate Setup windows, respectively.
- 2. Selected Protocol area: Located in the bottom left of the Setup window, the Selected Protocol area displays the details of the protocol selected in the File Browser area. The selected protocol file name appears at the top of this window. If you have selected a data file, the iQ5 software displays the protocol used to create the data file and the data file name. The iQ5 software the selected protocol both graphically and in spreadsheet format in the Selected Protocol area.
- 3. Selected Plate Setup area: Located in the bottom right of the Setup window, the Selected Plate Setup window displays the details of the plate setup selected in the File Browser area. The selected plate setup name appears at the top of this window. If you have selected a data file, then the plate setup within the data file and the data file name are both selected in the iQ5 software. When you select the Data File tab and a data file, the iQ5 software displays the Original and Current plate setup. The Original plate setup is the plate setup used to create the data file. The Current plate setup is the plate setup.
- 4. **Selected Data File area**: Located in the top right of the Setup window, the Selected Data File area displays the selected data file or the run set file name that you selected in the File Browser area. The Notes box displays the notes associated with either the run set or data file. Buttons in this area include:
 - Run: Used to initiate a real time PCR experimental run.
 - Begin End Point: Used to initiate an End Point run.
 - Analyze: Used to open a data file. You can also open the data file by double-clicking on the data file name in the File Browser area.

3.2 Plate Summary Window

The Plate Summary window provides a more detailed view of the plate including standard quantities and identifiers, as shown below. It is useful to check the Plate Setup after editing or creating the Plate and also before performing a run. It can also be used to check the plate layout in a Data File.

If you select the Data File tab, the plate shown in the Plate Summary window depends on whether you selected Original or Current in the Selected Plate Setup window. Original is the plate setup that the data file was created with. Current is the last saved plate setup.

	rwing Plate:	Sample	Plate.pts		Expe	eriment	Mc koperno	17 - T	1	Copy To (Clipboard	Pr	int
0	FAM)HEX	OTexasRe	d Ocy5									
	_	1	2	3	4	5	6	7	8	9	10	11	12
A	SampleType		Pos Ctrl-1		Unkn-1	Unkn-2	Unkm-3	Unkn-4	Unkn-5	Unico-6		Neg Ctrl-1	NTC-I
	copy number		N/A		N/A	N/A	N/A	N/A	N/A	N/A		N/A	N/A
1	Identifier		in the second									-RT	1 Calling
	SampleType	1	Pos Ctrl-1		Unkn-1	Unkn-2	Unico-3	Unkn-4	Unkn-5	Unkn-6	9	Neg Ctrl-1	NTC-I
в	copy number		N/A		N/A	N/A	N/A	N/A	N/A	N/A		N/A	N/A
	Identifier											-RT	
	SampleType		Pes Ctrl-1		Unkn-1	Unkn-2	Unkm-3	Unkn-4	Unkn-S	Unkn-6	5	Neg Ctrl-1	NTC-I
C	copy number		N/A		N/A	N/A	N/A	N/A	N/A	N/A	1	N/A	N/A
	Identifier											-RT	
	SampleType							S					
D	Concentration												
	Identifier	-	-					-					
	SampleType												
ε	Concentration												
	Identifier	-	-					-					
	SampleType			Std-1	Std-2	Std-3	Std-4	Std-5	Std-6	Std-7	Std-8		
F	copy number	-		1.00e+08	1.00e+07	1.00e+06	1.00e+05	1.00e+04	1.00e+03	1.00e+02	1.00e+01		
	Identifier	-	-	-			-	-					_
	SampleType	-		Std-1	5td-2	Std-3	Std-4	Std-5	Std-6	Std-7	5td-8		
6	copy number			1.00e+88	1.00e+07	1.00e+06	1.00e+05	1.00e+04	1.00e+03	1.00e+02	1.00e+01		
	Identifier	-	-	-		52d-3	Std-4			Std-7	-		-
	SampleType	-		Std-1	Std-2	Statement of Statement and	al and a second second second	Std-5	Std-6		Std-8		
н	copy number Identifier			1.00e+08	1.00e+07	1.00e+06	1.008+05	1.000+04	1.00e+03	1.00e+02	1.00e+01		

Fig. 3.2. The Plate Summary tab.

The spreadsheet displays data one dye layer at a time, and you can use the fluorophore buttons to select the plate summary of the desired dye layer, which is especially useful in Per Dye Layer edited plates.

To print the selected plate setup display, click Print. To copy and paste the entire plate setup, click Copy to Clipboard. You can also copy only a portion of the plate setup by clicking and dragging your mouse over the desired wells, then pressing CTRL+C in the iQ5 software, and then pressing CTRL+V in the desired application.

3.3 Plate Setup

The iQ5 and MyiQ real time PCR detection systems monitor only the wells defined to have some type of sample and at least one fluorophore. In the Plate Setup Editor, you specify the type of sample in each well and the fluorophores present in each well. If you define standards on the plate, the Plate Setup Editor can calculate the concentration of each standard in a dilution series given the appropriate information, or you can enter them manually. You can import a unique Identifier for each well from a CSV file or enter them manually. You can get to the Plate Setup Editor from the main Workshop module window.

3.3.1 Selecting a Plate Setup

To select an existing Plate Setup File in the Workshop module:

1. Click Plate and negotiate the directory to find the desired plate setup file. The vertical gray bar next to the directory tree can be dragged to the right to widen the window.

Note: A plate setup within a Data File can also be selected and viewed

2. Click on the file name and the plate setup will be displayed in the bottom right pane.

The sample type is displayed for every well one dye layer at a time. The buttons over the plate are used to select the currently displayed fluorophore. You can also determine what other fluorophores are in each well, though you cannot tell what the sample type is in those other dye layers.

The experimental sample volume as well as the type of vessel and sealing mechanism are displayed above the plate as shown below.

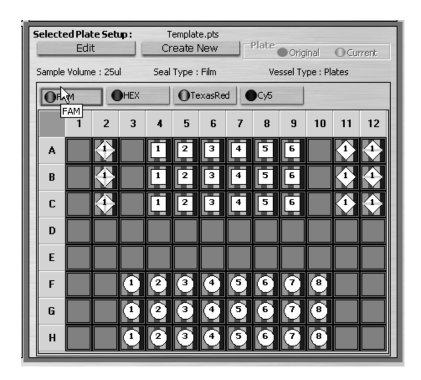


Fig. 3.3. The Selected Plate Setup information.

If you are looking at a data file that contains a plate setup that was edited and saved after the experimental data were collected, you can revert to the original plate setup by clicking Original in the Plate box. To go back to the present definition, click Current.

None of the information displayed on the plate in the Workshop home window can be edited from this screen.

3.3.2 Editing or Creating a Plate Setup

Editing a Plate Setup File

- 1. Select a Plate Setup
- 2. Click Plate or Data File and negotiate the directory to find the desired plate setup file. The vertical gray bar next to the directory tree can be dragged to the right to widen the window.
- 3. Click on the file name and the plate setup will be displayed in the bottom right pane.
- 4. Click Edit to bring the plate setup into the Plate Setup Editor.

Alternatively, you can double click on the file name in the directory and bring the plate setup directly into the Plate Setup Editor.

- 5. Use the Plate Setup Editor to edit the Plate, then click Save & Exit Plate Editing.
- 6. Confirm that the Plate is defined correctly using the Plate Summary.

Creating a New Plate Setup File

1. Click Create New in the bottom right pane of the home screen in the Workshop. It does not matter what plate setup is displayed at the time.

- 2. The software will transfer to the Plate Setup Editor.
- 3. Use the Plate Setup Editor to edit the Plate, then click Save & Exit Plate Editing.
- 4. Confirm that the Plate is defined correctly using the Plate Summary.

3.4 Plate Setup Editor

The Plate Setup Editor is comprised of a 96-well plate layout, functions for specifying the sample type and fluorophores in each well, and a spreadsheet displaying the definition in each dye layer for any individual well.

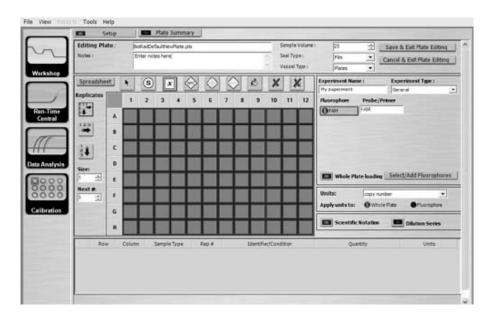


Fig. 3.4. The Plate Setup Editor.

There are two other windows accessible from the Plate Setup Editor:

- 1. Plate Summary shows well definition one dye layer at a time
- 2. Plate Setup Editor Spreadsheet is a comprehensive set data for all 96 wells at once, shown one dye layer at a time.

Exiting the Plate Setup Editor

You can return to the Workshop at any time by pressing Cancel & Exit Plate Editing, though you will lose your current work.

To save your work before returning to the home screen, click Save & Exit Plate Editing and assign a name to your new plate setup in a standard Windows Save dialog box. All plate setup files are automatically assigned an extension of .pts.

Note: Plate setup files from previous versions of the iQ or MyiQ software can be opened in the new software, but before it can be saved you must assign a vessel type and sealing mechanism.

3.4.1 Well Definition Icons



Fig. 3.5. The Well Definition Toolbar.

There are nine icon buttons running across the top of the representation of the experimental plate. These buttons are used to provide the two pieces of information required for each well: sample type and fluorophore(s) to be monitored. The active button is always surrounded by a red box.



Standard Pointer: used to select - but not alter wells.



Standard: Defines Standards



Unknown: Defines Unknowns



No Template Control: Defines No Template Controls



Positive Control: Defines Positive Controls



Paint Bucket Icon: When the Paint Bucket icon is active and you click a well in the plate setup, all wells in that replicate group will be assigned the currently selected fluorophore.



Delete Fluorophore Icon: When the Delete Fluorophore Icon is active and you click in a well in the plate setup, the currently selected fluorophore is removed from all wells in that replicate group. It does not change the definition of the sample type in that well.



Delete All Icon: When the Delete All icon is active and you click a well, all information from that well is removed from that well. The other members of a replicate group are not affected.

3.4.2 Editing a Plate Setup

- 1. From the Workshop module:
 - a. Click Create New in the plate setup display window to enter the Plate Setup Editor.

or

b. Click Plate. Open the desired plate setup file in the appropriate directory. Double click the file name to go directly to the Plate Setup Editor.

or

c. Click Plate. Open the desired plate file in the appropriate directory. Click the file name to open the plate setup in the bottom right pane of the Workshop window. Click Edit to open the plate setup in the Plate Setup Editor.

or

- d. Click Data File. Open the desired data file in the appropriate directory. Click the file name to open the plate setup associated with the data file in the bottom right pane of the Workshop window. Click Edit to open the plate setup in the Plate Setup Editor.
- 2. Enter or edit any notes about the plate setup in the Notes box.
- 3. Enter or edit the Sample Volume, Seal Type, and Vessel Type.
- 4. Enter or edit a name for the experiment.
- 5. Click Select/Add Fluorophores and select or edit the fluorophores to be used on the plate. Associate each fluorophore with a unique color.
- 6. Clear the Whole Plate Loading check box if it is not applicable; otherwise, leave the check box selected. If you are editing a plate, the Whole Plate Loading check box may be unavailable because it is not appropriate based on the current definition of the plate.
- 7. Click a sample type icon.
- 8. Select the type of replicate loading desired.
- 9. Click a fluorophore.
- 10. Click or drag across the plate to define wells with the selected fluorophore and sample type.
- 11. Continue defining the remaining wells that will contain the first fluorophore by changing to the other sample type icons.
- 12. To delete a previously defined well, click the Delete All icon, and then click the well.
- 13. To delete the selected fluorophore from a previously defined well, click the Delete Fluorophore icon, and then click the well.
- 14. If there are standards in the first dye layer, click Dilution Series and calculate the concentrations of the standards, or type the concentrations in the spreadsheet manually.

FOR SINGLE-COLOR EXPERIMENTS:

15. Click Save & Exit Plate Editing. Give the plate setup file a new name or overwrite the current version of the file. The extension .pts will be added automatically.

FOR MULTI-COLOR EXPERIMENTS WITH WHOLE PLATE LOADING SELECTED:

- 15. Click the second fluorophore.
- 16. To delete the second fluorophore from a previously defined well, click the Delete Fluorophore icon and then click the well.
- 17. Click the Paint Can icon.
- 18. Click or drag across any previously defined wells that will also contain the second fluorophore.
- 19. Now use the other sample type icons as necessary to define any wells that will contain the second fluorophore, but not the first.
- 20. Repeat steps 15–19 for the third, fourth and fifth fluorophores.
- 21. Click Save & Exit Plate Editing. Give the plate setup file a new name or overwrite the current version of the file. The extension .pts will be added automatically.

FOR MULTI-COLOR EXPERIMENTS WITH WHOLE PLATE LOADING DESELECTED:

- 15. Click the second fluorophore. All wells defined previously are now shown colored by the first fluorophore, but without indication of sample type. When Whole Plate Loading is deselected, no assumption is made about the sample type in the second and subsequent dye layers. The sample type can vary from dye layer to dye layer.
- 16. To delete the second fluorophore from a previously defined well, click the Delete Fluorophore icon and then click the well.
- 17. Click a Sample Type icon.
- 18. Click or drag across the plate to define wells with the second fluorophore and selected sample type.
- 19. Continue defining the remaining wells that will contain the second fluorophore by changing to the other sample type icons.
- 20. If there are standards in the second dye layer, click Dilution Series and calculate the concentrations of the standards, or enter the concentrations in the spreadsheet manually.
- 21. Repeat steps 15-20 for the third, fourth and fifth fluorophores.
- 22. Click Save & Exit Plate Editing. Give the plate setup file a new name or overwrite the current version of the file. The extension .pts will be added automatically.

3.4.3 Fluorophore Selection

The iQ5 and MyiQ real time PCR detection systems monitor only the wells defined to have some type of sample and at least one fluorophore. When using the iQ5 system, you may specify as many as five different fluorophores on a single plate. In addition, all five fluorophores may be in a single well, provided that each fluorophore is detected through a different filter pair.

When using the MyiQ system, each monitored well must only contain a single fluorophore. All selected fluorophores must all be detectable through the MyiQ's single filter pair. Fluorophores specified to be read through the same filter pair are known as "filter mates". Up to five fluorophores may be designated on a single plate when using the MyiQ system, provided that all five fluorophores are filter mates.

For more information on filter mates, refer to the following section.

3.4.4 Filter Mates

A new feature in the iQ5 software is the ability to choose to monitor as many as five different sets of wells with the same filter pair (that is, in the same dye layer) as shown below. For example, you may have five different experiments on the same plate, all using SYBR Green, and each experiment will be treated independently. When more than one fluorophore is independently monitored by a single filter pair, those fluorophores are called filter mates.

Note: The software cannot distinguish between filter mates in the same well; therefore, in all cases, filter mates have to be in separate wells. For example, you may not put FAM and SYBR Green in the same well.

Experiment Nan	ne:	Experiment Type :					
My experiment		General					
Fluorophore	Probe/Pri	ner					
O FAM	FAM						
HEX	HEX						
O TexasRed	TexasRed						
●Cy5	Cy5						
Whole Pla	ate loading	Selec	:t/Add Fluoropl	nores			

Fig. 3.6. List of five sets of wells.

Click Select/Add Fluorophores to open the Fluor Selector box. From this box, choose up to five fluorophores to be monitored. To remove a fluor from the list to be monitored, uncheck the box next to its name.

To change the default color associated with a fluor, click the color box on the row with the fluor name and choose a new color as shown below.

Filter Position	Fluor	Selected	Color
2	FAM		Lime
	SYBR		
	SYBR1		
	SYBR2		
	SYBR3		
	SYBR4		
	SYBR5		
3	HEX		Green
	JOE		
	TET		
	VIC		
4	Cy3		
	TAMBA		
5	TexasRed		SkyBlue
or Name :			Correct
er Position :	2	-	Cancel

Fig. 3.7. The Fluor Selector window.

You may add new dyes to the list at any time by clicking Add New Fluor. When you add a new dye, you must specify the filter to be used to monitor that fluor and you must later perform a pure dye calibration for that fluor.

3.4.5 Whole Plate Loading

There are two pieces of information required for each well: sample type and fluorophore(s) to be monitored. There are two ways of providing that information to the plate setup editor. When Whole Plate Loading selected, you specify the sample type and fluorophores separately. This is the most efficient way of defining a plate setup for a multicolor experiment when all wells have the same sample type at each dye layer, for example, you might choose to monitor four fluors in well A1, and in each dye layer, the sample type is Unknown. Note that if you choose Whole Plate Loading, it is assumed that standards in each dye layer are at the same concentration, that is, the FAM standard might be at 500 copies and then, by definition, the Texas Red standard is also at 500 copies.



When the sample types vary within a well, or in the special case of standards, when the concentration of standard varies with the dye layer (monitored fluorophore), then deselect Whole Plate Loading. When Whole Plate Loading is deselected, you specify the type of sample and fluorophore simultaneously, one dye layer at a time.

3.4.6 Specifying Replicates

The buttons for specifying replicates run down the left side of the representation of the experimental plate.



This is the default setting and indicates that each sample is identical (i.e., a single replicate).

With this button active, if you:

• Click an unused well, the sample type is incremented by one.

- Click a row letter or column number, the sample type for every member of the row or column is assigned the identical replicate number.
- Drag across a selection, the sample type for every member within the selection is assigned the identical replicate number.



The Horizontal and Vertical Replication buttons, together with the number in the Size box, specify the direction and number of replicates to be automatically defined.

With the horizontal direction (row) button active and the Size set at 5, if you:

Click a well, the sample type is incremented by one for the next five wells in the horizontal direction, each of the five wells will

contain an identical replicate number. If there are not five wells left in the row, the software will wrap around to the next row.

Click a row letter, the sample number of the first five wells will be incremented by one, the next five wells
incremented by two, the last two wells will be incremented by three. The software does not automatically
wrap around so that the last set of
replicates has the same size as the others.

replicates has the same size as the others.

Drag across a selection, the first five members of the selection are incremented by one, the second five
members are incremented by two, etc. Numbering goes in a horizontal direction and wraps around to the
next row within the selection. The last set of replicates may have a smaller size than the others, depending
on the number of wells within the selected area

With the Vertical direction (column) button active and the Size set at 3, if you:

- Click a well, the sample type is incremented by one for the next three wells in the vertical direction. If there are not three wells left in the column, it will wrap around to the next column.
- Click a column number, the sample number of the first three wells will be incremented by one, the next three wells

incremented by two, the last two wells will be incremented by three. The software does not automatically wrap around so that the last set of replicates has the same size as the others.

 Drag across a selection, the first three members of the selection are incremented by one, the second three members are

incremented by two, etc. Numbering goes in a vertical direction and wraps around to the next column within the selection. The last set of replicates may have a smaller size than the others, depending on the number of wells within the selected area.



Size:

÷

The Next # box allows you to overwrite the default replicate number for the next sample to be defined. This is an easy way to correct a mistakenly assigned replicate number. Use the up and down arrows to select the next desired replicate number, or highlight the field and type the desired number.

3.4.7 Sample Volume, Seal Type, and Vessel Type

In Plate Setup you must specify the sample volume, the type of sealing and the type of vessel to be used in the experiment. This is to properly associate pure dye calibration files with the data file, since pure dye calibrations are unique to each specific volume, vessel and sealing combination. Use the up and down arrow keys to specify the sample volume, or highlight the field and type in the volume. Select the seal type and vessel type from the pull down menus.

Sample Volume :	25	÷	Save & Exit Plate Editing
Seal Type :	Film	•	Cancel & Exit Plate Editing
Vessel Type :	Plates	-	

Fig. 3.8. The Seal and Vessel Type Dialog Box.

The sample volume may be entered directly or input through the up and down arrows.

3.5 Dilution Series

The iQ5 software can automatically calculate the concentrations of each member of a dilution series, given the dilution factor and the concentration of either the most-concentrated or least-concentrated sample, in the Dilution Series box as shown below.

Scientific Notation	Dilution Series
Define Standards: Replicates from :	1 ÷ to: 8 ÷
Starting Concentration :	1.00E-03
Series Identifier :	
Dilution Factor :	10 Increasing Decreasing
Define Close	Apply Dilution Series

Fig. 3.9. The Dilution Series box.

3.5.1 Defining a Dilution Series of Standards

- 1. Define the range of standard Replicates to be defined. By default, the beginning and ending standards are the I owest- and highest-numbered standards, but you may define only a subset of all standards by clicking on the up or down arrows.
- 2. Enter the Starting Concentration, the concentration of the lowest-numbered standard to be defined. You may enter the concentration using scientific notation, even if the Scientific Notation button is unselected. The reverse is also true-you may make an entry *without* using scientific notation if the Scientific Notation button is selected. The Scientific Notation button controls only how the data are displayed in the spreadsheet.
- 3. Enter a Series Identifier for the standard if desired. It will be shown in the spreadsheet and in the reports.
- 4. Enter the fold Dilution Factor. The default value is 10.
- 5. Indicate whether the concentrations decrease or increase as the standard replicates increase in number. The default is Decreasing.
- 6. Click Apply Dilution Series and all standard concentrations will be automatically calculated.

3.5.2 Adding Additional Standards

When Define is selected (at the bottom of the dialog box), any standards added to the set after you close the Dilution Series box, will have concentrations automatically calculated.

If Define is not selected, then in order to assign values to any newly added standards, you must open the Dilution Series dialog box again, make any necessary edits and click Apply Dilution Series.

3.6 Plate Summary

The Plate Summary window provides a more detailed view of the plate including standard quantities and identifiers, as shown below. It is useful to check the Plate Setup after editing or creating the Plate and also before performing a run. It can also be used to check the plate layout in a Data File.

If you select the Data File tab, the plate shown in the Plate Summary window depends on whether you selected Original or Current in the Selected Plate Setup window. Original is the plate setup that the data file was created with. Current is the last saved plate setup.

Vi	ewing Plate:	Sample	Plate.pts		Expe	riment	N/ societte	- 50	1	Copy To Clipboard Print			
Q	FAM)HEX	OTexasRe	d Ocy6									
		1	2	3	4	5	6	7	8	9	10	11	12
	SampleType		Pos Ctrl-1		Unkn-1	Unkn-2	Union-3	Unkn-4	Unkn-5	Union-6		Neg Ctrl-1	NTC-1
A	copy number		N/A		N/A	N/A	N/A	N/A	N/A	N/A	1	N/A	N/A
	Identifier		and the second second				0.000					-RT	
	SampleType	-	Pos Ctrl-1		Unkn-1	Unkn-2	Unkm-3	Unkn-4	Unkn-5	Union-6		Neg Ctrl-1	NTC-1
B	copy number		N/A		N/A	N/A	N/A	N/A	N/A	N/A	-	N/A	N/A
1	Identifier	_										-RT	
	SampleType	· · · · ·	Pos Ctrl-1		Unkn-1	Unkn-2	Union-3	Unkn-4	Unkn-5	Union-6		Neg Ctrl-1	NTC-I
C	copy number		N/A		N/A	N/A	N/A	N/A	N/A	N/A		N/A	N/A
	Identifier											-RT	
	SampleType	-	-	-	1						-		
D	Concentration				-						-		
_	Identifier	-	_										
	SampleType	-							1				
E	Concentration												
	Identifier		-		and the second								_
	SampleType			Std-1	Std-2	Std-3	Std-4	Std-5	Std-6	Std-7	Std-8		
F	copy number			1.00e+08	1.00e+07	1.00e+06	1.00e+05	1.00e+04	1.00e+03	1.00e+02	1.00e+01		
	Identifier		-		1	-		-	-	1	-		
	SampleType			Std-1	Std-2	5td-3	5td-4	Std-5	5td-6	Std-7	5td-8		
G	copy number	-		1.00e+08	1.00e+07	1.00e+06	1.00e+05	1.00e+04	1.00e+03	1.00e+02	1.00e+01		
	Identifier			1		-			1	-	1		
	SampleType			Std-1	Std-2	Std-3	Std-4	504-5	Std-6	Std-7	51.0-8		
н	copy number			1.00e+88	1.00e+07	1.00e+06	1.00e+05	1.00c+04	1.00e+03	1.00e+02	1.00e+01		

Fig. 3.10. The Plate Summary tab.

The spreadsheet displays data one dye layer at a time, and you can use the fluorophore buttons to select the plate summary of the desired dye layer, which is especially useful in Per Dye Layer edited plates.

To print the selected plate setup display, click Print. To copy and paste the entire plate setup, click Copy to Clipboard. You can also copy only a portion of the plate setup by clicking and dragging your mouse over the desired wells, then pressing CTRL+C in the iQ5 software, and then pressing CTRL+V in the desired application.

3.6.1 Spreadsheets

There are two different spreadsheets available from the plate setup editor. One spreadsheet opens beneath the representation of the experimental plate any time that the pointer tool is clicked on a well. This spreadsheet simultaneously displays information about each dye layer in the active well.

In this spreadsheet you may edit the Sample Type, Identifier and Quantity. Any changes are carried out through all members of a replicate group. It is also possible in this spreadsheet to change the replicate group assignment of a well, but it will also automatically change the assignment of all other members of the original replicate group.

Note: In order to edit the replicate group assignment of a single well, without changing the other members of the same replicate group, make the change on the plate, not in the spreadsheet.

Row	Column	Sample Type F		Rep #	Identifier/Condition	Quantity	Units	
F	4 Standard		Standard 🖸 2		mouse cDNA	1.00E+07	copy number	
		Standard V Standard V		2	mouse cDNA	1.00E+07	copy number	
				2	mouse cDNA	1.00E+07	copy number	
		Standard		2	mouse cDNA	1.00E+07	copy number	

Fig. 3.11. The Well Identifier Spreadsheet.

3.6.2 Plate Setup Editor Spreadsheet

The second spreadsheet can be opened by clicking the Spreadsheet button in the plate setup editor window. This spreadsheet displays information about all wells simultaneously, one dye layer at a time. Dye layer selection is made at the top of the spreadsheet.

ate Spreadsl	noocearch to		FAM	Import Ic	EAILF	late Spreadsheet Editing
Row	Column	Sample Type	Rep #	Identifier/Condition	Quantity	Units
В	4	Unknown	1	0 Hrs	N/A	copy number
В	5	Unknown	2	1 Hr	N/A	copy number
В	6	Unknown	3	2 Hrs	N/A	copy number
С	4	Unknown	1	0 Hrs	N/A	copy number
С	5	Unknown	2	1 Hr	N/A	copy number
С	6	Unknown	3	2 Hrs	N/A	copy number
D	4	Unknown	1	0 Hrs	N/A	copy number
D	5	Unknown	2	1 Hr	N/A	copy number
D	6	Unknown	3	2 Hrs	N/A	copy number
F	3	Standard	1		1.00E+08	copy number

Fig. 3.12. The Plate Editor Spreadsheet.

Within this spreadsheet you may change the Sample Type, the Identifier, and the Quantity. Note that changes made to one member of a replicate group are carried through to all members of the replicate group. Replicate group assignment changes should be made on the plate, not in the spreadsheets.

- In Whole Plate Loading, changes made to any dye layer within a well are extended to the other dye layers within the well upon pressing Enter or clicking in another cell of the spreadsheet. These changes are then extended to every other well within the replicate group.
- When Whole Plate Loading is toggled off, changes made to one dye layer are considered unique to the dye layer and are not automatically extended to the other dye layers in the well. However, these changes are extended in the same dye layer to every other well within the replicate group.

The single-dye layer spreadsheet has another feature that differentiates it from the single-well spreadsheet. From this spreadsheet you may Import Identifiers from an external CSV (comma separated values) file. The number of identifiers must match the number of wells before the import can be carried out. The simplest way to use the import feature is to fill out the Identifier template file provided with the software and then save the file in the CSV format. The Identifier template file is called "Well Sample Identifier Import.xls, and is located in the User1 folder of the iQ5 SW directory.

3.7 Protocol

The iQ5 software allows you to create and run thermal cycling programs in the iCycler thermal cycler and to simultaneously collect and analyze fluorescent data captured by the iQ5 or MyiQ optical module. Customized files, called Protocols, direct the operation of the iCycler and also specify when optical data will be collected during the thermal cycling run. iQ5 protocol files are stored with the extension .tmo.

Protocol files contain the information necessary to direct the operation of the iCycler and also instruct the iQ5 or MyiQ camera when to collect data.

- A protocol is made up of as many as 9 cycles and a cycle is made up of as many as 9 steps.
- A step is defined by specifying a setpoint temperature and the dwell time at that temperature.
- A cycle is a collection of steps that are repeated (up to 600 times).

Every protocol must have at least one data collection step. This may be a real-time data collection step or a melt curve data collection step.

3.7.1 Protocol Viewer

The Protocol Viewer is shown below. It is selected by clicking on the Protocol button. When active, a green border appears around the Graphical Display and the spreadsheet.

Protocol	E F	Plate	🗖 Run	Set	🔲 Data File					
Drives Drives Drives Step StepAmp+Mek Shotcut to Code StepAmp+Mek Shotcut to Code Masks Persisten ReportTe ReportTe ReportTe ReportTe SampleFi One Step RT-PCR One Step RT-PCR SampleFi One Step RT-PCR+Met SampleFi Codecade Protocol: FourColor.tmo within Sample Gene Expression - Multiple										
Selected Protocol : Edit	F Create		o within Sam Protocol	ole Gen						
Cycle 1		Cycle	2 50X							
Step 1		Step	1		Step 2					
95.0		95.0			55.0					
3:00		0:30			1:00					
					Ô					
Cycle Repeats	Step	Dwell Time	Setpoint	PC	R / Melt Data Acquisition					
1 1										
	1	3:00	95.0							
2 50										
	1	0:30	95.0							
	2	1:00	55.0	Ô	Real Time					
	-									

Fig. 3.13. The Protocol Viewer.

When the Protocol button is selected (as indicated by the green active button) the browser displays protocol files (which have the suffix .tmo)

The browser may be used to select a protocol which is displayed in both the Graphical Display and the Spreadsheet.

The size of the folder browser and the file pane can be changed by moving the bar that separates them.

Temperature and Dwell Time Ranges

- Temperatures between 4.0°C and 100.0°C may be entered for any set point temperature.
- Finite dwell times may be as short as 1 second (00:01) or as long as 99 minutes and 59 seconds (99:59).

Thermal protocols can be selected, created and edited from the Protocol tab in the Workshop module.

3.7.2 Selecting a Protocol

- Use the browser below the Protocol button to locate the folder the Protocol is located in, and then click on the protocol name to select it.
- The selected protocol appears in the Selected Protocolwindow.
- The iQ5 software has a number of sample protocol files you can use.

3.7.3 Editing or Creating a Protocol

To edit a selected protocol, double click the protocol file or click Edit in the Selected Protocol window.

To create a protocol from a protocol template, click Create in the Selected Protocol window.

Clicking Edit or Create in the Viewing Protocol window will transfer the software to the Protocol Editor. You can edit the protocol in the spreadsheet at the bottom of the Editing Protocol window.

Editing Dwell Time and Setpoint Temperature

Click in the spreadsheet in the Dwell Time or Setpoint cell, then enter the Dwell Time or Setpoint temperature.

Data Acquisition

Click in the spreadsheet in the Data Acquisition column and select Real-Time data collection for any step.

Note: Ensure that every thermal protocol has at least one Data Acquisition step.

Chose Melt Curve if data from a melt curve is required.

Inserting Cycles and Steps

To insert a cycle click in the Insert column on the Cycle row. Cycles are indicated with a blue background.

To insert a step click in the Insert column on a Step row. Steps are indicated with a white background.

Deleting Cycles and Steps

To delete a cycle click in the Delete column on the Cycle row. Cycles are indicated with a blue background.

To delete a step click in the Delete column on a Step row. Steps are indicated with a white background.

Add a Melt Curve Step

Add a Gradient Step

Saving the Protocol

To save a protocol Click Save & Exit Protocol Editing. Enter the name of the protocol in the Save As dialog box and click Save again.

The Protocol Editor may only be exited by clicking **Save & Exit Protocol Editing** or **Cancel & Exit Protocol Editing**.

3.7.4 Add Protocol Options

If you want to add any protocol options, first enable them by clicking in the check box next to its description in the Show Options box. The following options are available:

- 1. Gradient
- 2. Infinite Hold
- 3. Ramping
- 4. Temperature Change
- 5. Time Change
- 6. Cycle Description
- 7. Step Process

Infinite Hold

When a cycle is not repeated, you can specify the dwell time at any step in that cycle as infinite by using the Infinite Hold option. This means that the instrument will maintain the specified temperature until execution is interrupted. When an infinite dwell time is programmed within a protocol at some step other than the last step, the instrument will go into Pause mode when it reaches that step and will hold that setpoint temperature until the Continue Running Protocol button in the Thermal Cycler tab of the Run-Time Central Module is selected.

An infinite hold may be programmed in the following way:

1. Click Infinite Hold in the Show Options box. A new column titled Hold appears in the spreadsheet.

2. Select the Hold check box for the step that you want to maintain at a constant temperature and enter the desired temperature in the Setpoint cell of the spreadsheet.

Ramping

The ramp rate is the speed with which the iCycler changes temperatures between the steps of a cycle, or between cycles. The default condition is for the iCycler to adjust temperatures at the maximum ramp rate. The iCycler allows you to change temperatures at a fixed rate less than the maximum.

Ramp rates are adjustable to 0.1°C/sec and must fall within the range of 0.1 to 3.3°C per second for heating and 0.1 to 2.0°C per second for cooling. Invalid ramp rate entries are adjusted to the nearest valid entry.

A ramp rate may be programmed in the following manner:

- 1. Click Ramping in the Show Options box. A new column titled Ramp Rate will appear in the spreadsheet.
- Double click in the Ramp Rate column on the line of the spreadsheet containing the temperature toward which you
 wish to control the ramp rate. Use the pull down menu to select MIN or MAX or make a direct entry into the field. If
 an invalid ramp rate is input, it is adjusted to the nearest valid ramp rate automatically.

Temperature Change

You may program an automatic periodic increase or decrease in the step temperature in a repeated cycle. Temperature increments or decrements may be as little as 0.1°C per cycle. You may make the increase or decrease as frequently as every cycle, and the increase or decrease can begin following any cycle. The temperature increment or decrement may be as large as desired, as long it does not result in temperatures which are outside the temperature limits of 4°–100°C.

A temperature increment or decrement may be programmed in the following way:

- 1. Click Temperature Change in the Show Options box. Three new columns will appear in the spreadsheet.
- 2. For the repeated step you want to affect, enter the incremental change desired in the Temperature change column. To decrement the temperature enter the decremental change as a negative number (eg -0.5).
- 3. Enter the repeat in which you want the change to occur for the first time in the Begin Repeat column. Usually it is cycle 2, but it can be any cycle greater than 1.
- 4. Enter the frequency that you want the change to occur in the How Often column. Usually you will want the change to occur every repeat, so enter 1 in this column.

Time Change

You may program an automatic periodic increase or decrease in the step dwell time in a repeated cycle. Time increments or decrements may be as little as 1 sec per cycle. You may make the increase or decrease as frequently as every cycle, and the increase or decrease can begin following any cycle. The time increment or decrement may be as large as desired, as long it does not result in dwell times which are outside the limits of 00:00 and 99:59.

A time increment or decrement may be programmed in the following way:

- 1. Click Time Change in the Show Options box. Three new columns will appear in the spreadsheet.
- 2. For the repeated step that you want to affect, enter the change desired in the Time Change column. To decrement the time enter the decremental change as a negative number (e.g., -0:05).
- 3. Enter the cycle in which you want the change to occur for the first time in the Begin Repeat column. Usually it is cycle 2, but it can be any cycle greater than 1.
- 4. Enter the frequency that you want the change to occur in the How Often? column. Usually you will want the change to occur every repeat, so enter 1 in this column.

Cycle Description/Step Process

You can choose from a list of descriptive names or enter one of you own to describe cycle or step processes, respectively.

- A cycle description or step process may be entered in the spreadsheet in the following manner:
- 1. Click Cycle Description or Step Process in the Show Options box.
- 2. Click the cell of the spreadsheet for the cycle or step you wish to name and either choose one of the listed names from the pull down menu or enter one of your own.

3.7.5 Gradient

A thermal gradient may be programmed across the reaction block at any step of a protocol. The gradient runs from the back of the instrument to the front, with the hottest temperature in Row A and the coolest temperature in Row H. All wells in each respective row are at the same temperature so at any time during a gradient step, there will be eight different temperatures across the block with 12 wells at each different temperature. The gradient may be as large as 25°C or as small as 1°C. The gradient is not linear, but is highly reproducible. No row can be at a temperature higher than 100°C or lower than 40°C during the gradient step.

The gradient may be programmed in the following way:

- 1. Click Gradient in the Show Options box. Two columns will appear in the spreadsheet and a representation of the gradient will appear on the right side of the window.
- 2. Click the Gradient checkbox in the spreadsheet for the desired step.

Cycle	Repeats	Step	Dwell Time	Setpoint	PCR / Melt Data Acquisition				Range
1	1								
		1	3:00	95.0			-		
2	40								
		1	0:10	95.0			•		
		2	0:30	55.0	Ô	Real Time	•		10.0

Fig. 3.14. The Protocol Editing Table.

3. The temperature listed in the Setpoint cell of the spreadsheet will be the coolest temperature on the block during the gradient step (Row H). Enter the desired difference between the coolest and hottest temperatures during the gradient step in the Range cell of the spreadsheet. The Gradient Display will update with the temperatures at each row.

Alternative Gradient Programming Method:

Cycle Step									
Α	65.0								
В	64.5								
C	63.3								
D	61.4								
E	58.9								
F	57.1								
G	55.8								
н	55.0								
Rang	Range								
10.0									

If you want to obtain a specific temperature at any one row:

- 1. Click Gradient in the Show Options box. Two columns will appear in the spreadsheet and a representation of the gradient will appear on the right side of the window.
- 2. Click the Gradient checkbox in the spreadsheet for the desired step.
- 3. Enter the desired temperature into that row on the gradient display.
- 4. Press Enter.
- 5. The temperatures for the other rows will be calculated based on the input desired temperature and the range. You cannot specify the exact temperature on more than one row at a time.

Fig. 3.15. Gradient Display Table.

You can change the range in the spreadsheet or you can make a direct entry of the range in the gradient display. Press Enter and the display will update with the new calculated temperature for each row.

Note: A gradient can not be applied to any step which also has a Melt Curve

3.7.6 Melt Curve/Peak

Melt Curve/Peak analysis is a dynamic tool used to measure the melting temperature (T_m) of double stranded DNA molecules. DNA duplexes can be visualized by either incorporation of DNA-binding dyes (e.g. SYBR Green I) or by hybridization with fluorescently labeled probes. In the case of DNA-binding dyes and non-cleavable hybridization probes, fluorescence is brightest when the two strands of DNA are annealed. As the temperature is raised towards the T_m of the duplex, the fluorescence will decrease at a constant rate (constant slope). At the T_m , there is a dramatic reduction in the fluorescence with a noticeable change in slope. The rate of this change is determined by plotting the negative first derivative (-dF/dT) versus temperature. The greatest rate changes yield visible peaks, representing the T_m of the double-stranded DNA complexes.

Three major applications for melt curve analysis are:

- 1. Peak identification (number of amplified products)
- 2. Characterization of molecular beacons and
- 3. Allelic discrimination.

The first two applications are typically used as a guide for improving real-time PCR assay development. The third, allelic discrimination, is any assay used to distinguish or detect mutations between sequences of DNA. Single Nucleotide Polymorphism (SNP) Identification is one type of allelic discrimination assay. One way in which SNP detection can be done is to use the melt curve functionality in this software with fluorescent resonance energy transfer (FRET) assays.

A Melt Curve cycle may follow an amplification cycle or can be conducted independently of the amplification. The melt curve may be programmed in the following manner:

- 1. Melt curves are programmed as a repeated cycle containing only one step. The temperature is programmed to increase or decrease incrementally with each repeat of the cycle. The increase or decrease combined with the number of repeats may not result in a emperature that is below 4°C or above 100°C at any time during the protocol.
- 2. Insert a cycle into the protocol at the point that you want the melt curve. The step generated in this cycle will be used to generate the melt curve data.
- 3. Enter the temperature at which you wish to begin the Melt Curve in the Setpoint cell (4°C-100°C).
- 4. Enter an appropriate dwell time for data collection under the Dwell Time column. Dwell times for melt curve will vary based on the number of fluorophores in the experiment. To enter 10 seconds, type 0 followed by : then 10 (that is, as the time appears in the spreadsheet). Alternatively 10 seconds can be entered as 0.10.
- 5. Click in the PCR/Melt Data Acquisition column and select Melt Curve. A green camera will appear in the Data Acquisition cell as will add two additional columns to the spreadsheet entitled Temperature Change and End Temperature. By default Temperature Change is 0.5°C and the End Temperature is 95°C (unless the setpoint is 95°C). The number of repeats to achieve this melt curve is automatically calculated. The Temperature Change can be as low as 0.1°C increments. Typical Temperature Change values are 0.3–0.5°C for SYBR Green I.

Cycle	Repeats	Step	Dwell Time	Setpoint	PCR / Melt Data Acquisition			Temperature Change	End Temperature
1	1								
		1	1:00	95.0			•		
2	1								
		1	1:00	55.0			-		
3	81								
		1	0:10	55.0	ø	Melt Curve	Ŧ	0.5	95.0

Fig. 3.16. Melt Curve Protocol Editing.

- 6. Anneal Curves can be also created by entering the Temperature Change as a negative value eg -0.5
- 7. Save the protocol by entering a file name and then clicking Save & Exit Protocol Editing.

The minimum dwell times necessary for data collection are:

- 1 fluorophore = 10 sec
- 2 fluorophores = 20 sec
- 3 fluorophores = 30 sec
- 4 fluorophores = 40 sec
- 5 fluorophores = 50 sec

We recommend using a slightly higher dwell time than the minimum values so that more data points are collected at each repeat.

3.7.7 Sample Protocol Files

2Step.tmo: This protocol can be used for most single fluorophore or SYBR Green I real-time PCR experiments.

Cycle	Repeats	Step	Dwell Time	Setpoint	PC	R / Melt Data Acquisition
1	1					
		1	3:00	95.0		
2	40					
		1	0:10	95.0		
		2	0:30	55.0	Ô	Real Time

2StepAmp+Melt.tmo: This protocol can be used for most SYBR Green I real-time PCR experiments. The melt curve that follows the qPCR experiment can be used to determine the number of amplicons produced in the qPCR reaction

Cycle	Repeats	Step	Dwell Time	Setpoint	PC	PCR / Melt Data 1 Acquisition		Temperature Change	End Temperature	Begin Repeat	How Often?
4	1										
		1	3:00	95.0			Ŧ				
2	40						_				
		1	0:10	95.0			•				
		2	0:30	55.0	Ô	Real Time	•				
3	1						_				
		1	1:00	95.0			Ŧ				
4	1						_				
		1	1:00	55.0			•				
5	81										
		1	0:30	55.0	Ô	Melt Curve	Ŧ	0.5	95.0		

Gradient.tmo: This protocol can be used to determine the most optimal temperature to perform real-time PCR experiments.

Cycle Step	
Α	65.0
В	64.5
C	63.3
D	61.4
E	58.9
F	57.1
G	55.8
н	55.0
Ran	ge
10.0	

Cycle	Repeats	Step	Dwell Time	Setpoint		/ Melt Data		Gradient	Range
1	1								
		1	3:00	95.0			•		
2	40								
		1	0:10	95.0			•		
		2	0:30	55.0	Kõn R	eal Time	•	•	10.0

3.8 Run Set

A Run Set is a linked Plate Setup and Protocol and Run Set files generated by the iQ5 software are stored with the extension '.run'. Run sets are useful for experiments that are repeated on a frequent basis.

OUse Persistent Well Factors																	
Collect Well Factors from Experimental Hote							ning us	er sele	cted Pi	otocc	il and	Plate		-	Br	iqin R	un
ected	Protocal:		2Step.tm	,			ed Plate			DemoPf							
	Cycle 1			(C)	rde 2.40X	Sample	Volume			l Type			Ve	ssel Ty	pe:Pla	ites	_
	Step 1 95.0 °C			tep 1 5.0 °C	Step 2	Of/	м	01	EX	0	exasRo	đ					
-	3:00	_		5.0°C	60.0 °C 0:30		1	2	3 4	5	6	7	8	9	10	11	1;
					¢3				1								
Γ						B		0 0	0	0				-			-
					2	c		8 6	0	0							
						D		T	T	T							
Cycle	Repeats	Step	Dwell Tirre	Setpoint	PCR / Melt Data Acquisition	E		30	10	1						-	
1	1					F					123					181	
1.00	Theory and	1	3:00	95.0			-	-1-	4		-	-		-	-		-
2	40		0:30	95.0		6											
		1	0.30	73.0		H	1000				1000					1000	

Fig. 3.17. The Run Set Selection Window.

When the Run Set Tab is selected, run set files are shown in the File Browser.

A Run Set is a linked protocol and plate set up and run set files have the extension '.run'.

The protocol defined in the run set is shown in the Selected Protocol window and the plate setup defined in the run set is shown in the Selected Plate Setup window.

3.8.1 Selecting a Run Set

- Click Run Set.
- Use the browser below the Run Set button to locate the folder the run set file is located in and then click on the run set file name to select it.
- The selected run set will appear in the Selected Protocol and Selected Plate Setup windows.
- See an example Run Set.

3.8.2 Creating a Run Set

To create a run set, first select the plate setup and protocol that are to be linked.

The plate setup and protocol can be stand alone files (plate setup files have the extension .pts while protocol files have the extension .tmo) or they can be already together in a data file (with the extension .opd). A data file has two plate setups: Original is the plate set up the data file was created with, and Current is the plate setup the data file was last saved with. The run set will be created with the plate setup displayed in the Selected Plate Setup window.

Once you have chosen the protocol and plate setup files, click Create Run Set from the File menu.

3.9 Data File

Primarily used to select and open a data file. Can also be used to run a real-time experiment using the same protocol and plate setup that where used to create the data file. The plate setup that is selected can be either the Original plate setup or the Current (last saved) plate setup.

3.9.1 Selecting a Data File

- Click the Data File button.
- Use the file browser below the Data File button to locate the folder the data file is located in and then click on the data file name to select it.
- The protocol and plate setup used in the data file will appear in the Selected Protocol and Selected Plate Setup windows. The plate setup that is shown can be either the Original plate setup or the Current (last saved) plate setup. If the data file protocol and plate setup are being used as the conditions for running a new experiment, then the plate setup that is shown in Selected Plate Setup window will be the one that is used to run the new experiment.

3.9.2 Opening a Data File

To open an optical data file from an amplification or melt curve experiment, select the name of the data file in the File Browser and click Analyze (or double click the file name).

Opening a pure dye calibration file will not only show the pure dye fluorescent data in the PCR Quant screen but will also write this calibration file to the appropriate calibration folder. This will overwrite an calibration file of the same volume, reaction vessel, seal, optical camera and thermocycler and all subsequent experimental data will be collected and analyzed with the new calibration file.

3.9.3 Applying Alternate RMEs – iQ5 Systems Only

Optical data is saved to the data file along with the Calibration Files. It is possible to overwrite the calibration files stored within the opd data file thus changing the analysis of the experiment. This feature protects you from losing valuable experimental data if the calibrations were conducted incorrectly.

To apply an alternate RME (pure dye calibration file) to a dataset, you must first select the data file in the File Browser. Then select Apply Alternate Well Factors from the Tools menu. Use the Select Well Factors File Name dialog box to navigate to, then open the Well Factors calibration file. After the Well Factors calibration file has been opened, a Save Optical Data File dialog box will appear. The existing well factor calibration for this data file will be overwritten. For this reason it is recommended to save the optical data file with a different name from the original to ensure that the original file, with the original calibration data, is maintained for reference.

Do not use the well factor file containing dynamic well factors (identified by the name Dynamic_vessel_seal_volume_iQ5CameraSerial#_iCyclerSerial#) since this file contains only the well factors for the wells that were used in the experiment that generated this file. If the well list does not match, the operation will fail. Hence, it is a good practice to only use the Persistent Well Factor files when applying alternate well factors.

3.9.4 Applying Alternate Well Factors

To apply alternate Well Factors to a data file first select the data file in the File Browser. Then select Apply Alternate Well Factors from the Tools menu. Use the Select Well Factors File Name dialog box to navigate to, then open the Well Factors calibration file. After the Well Factors calibration file has been opened a Save Optical Data File dialog box will appear. The existing well factor calibration for this data file will be overwritten. For this reason it is recommended to save the optical data file with a different name from the original to ensure that the original file, with the original calibration data, is maintained for reference.

Do not use the well factor file containing dynamic well factors (identified by the name

Dynamic_vessel_seal_volume_iQ5CameraSerial#_iCyclerSerial#) since this file contains only the well factors for the wells that were used in the experiment that generated this file. If the well list does not match, the operation will fail. Hence, it is a good practice to only use the Persistent Well Factor files when applying alternate well factors.

Section 4 Run-Time Central Module

This module is entered automatically after clicking Run or Run End Point in the Workshop module. There are three windows in this module: Initiate Run, Show Plate and Monitor Run.

4.1 Initiate Run Window

Use the Initiate Run window, shown below, to confirm run conditions and then to initiate the optical data run. The protocol to be run is in the bottom left corner of the window, and the plate setup that will be used appears in the bottom right corner. The type of well factors to be used in the run can be selected in the top left section of the window. You can record details of the experiment in the Notes box, and these notes will be incorporated into the experimental file.

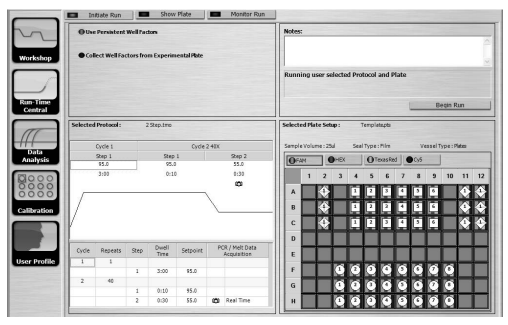


Fig. 4.1. The Initiate Run window.

To begin a run, click Begin Run. The Save dialog box opens. Type a name for the optical data file. The iQ5 software saves data automatically during the experiment.

4.1.1 Initiate Run Window - End Point Run

Initiate an End Point run by selecting a Plate Setup, then clicking Run End Point. The software transfers to the Initiate Run window in the Run-Time Central module. An End Point run uses a canned protocol in which only the temperature of the data collection step can be modified. Enter the desired temperature of the data collection step into the Setpoint: box as shown below. Click Enter.

Notes:		2
Running End Plate	Point Protocol and user selec	cted
		Begin Run

Fig. 4.2. The Endpoint Setpoint box.

Changes entered into the Setpoint: box appear in the Selected Protocol region of the Initiate Run window as shown below.

Cycle 1 2X	
Step 1	
Step 1 60.0 °C	
0:30 ເວັນ	
6	

Fig. 4.3. The Selected Protocol region of the Initiate Run window.

To begin a run, click Begin Run. The Save dialog box opens. Type a name for the optical data file. The iQ5 software saves data automatically during the experiment.

4.2 Show Plate Window

The Show Plate window can be used to visualize samples loaded into the reaction block of the iQ5 system. Use the Show Plate window to verify the position and orientation of samples prior to starting a run in the iQ5 or MyiQ instruments.

emplate.pts	Fluor Selection	Exposure Time (ms)
	• HEX	Expose
	TexasRed	Expose
	• Cy5	
		Show loaded wells
	Well Average Inner Fluoresc	All 0000.0
nstructions:		
 Click the Expose button to collect an exposure. An image of the sample area luorescence will be displayed above. Increase exposure time and collect a new 		
instructions: 1) Click the Expose button to collect an exposure. An image of the sample area luorescence will be displayed above. Increase exposure time and collect a new exposure if the fluorescent signal from the loaded wells is too low. 2) Confirm that small green boxes are visible around each well where sample luorescence will need to be detected.		

Fig. 4.5. The Show Plate window.

The following steps describe typical use of the Show Plate window:

- 1) Collect an exposure from the camera by clicking on the Expose button, an image of the fluorescence from the sample block area will be displayed.
 - a. **NOTE**: If the image displayed is too bright or too dim, it may be necessary to adjust the exposure time using the pull-down box in the right hand corner of the Show Plate window.
- 2) Confirm that the small green boxes are visible around each well where sample fluorescence will need to be detected.
 - a. If necessary, open the instrument and correct sample positions to match the plate setup for sample detection.
- 3) If a multiplex assay is being performed, repeat steps 1 & 2 for all fluors present in Fluor Selection list.
- 4) When finished, return to the Initiate Run screen

4.3 Monitor Run Window

Open the Monitor Run window by clicking Monitor Run. The window appears as shown below.

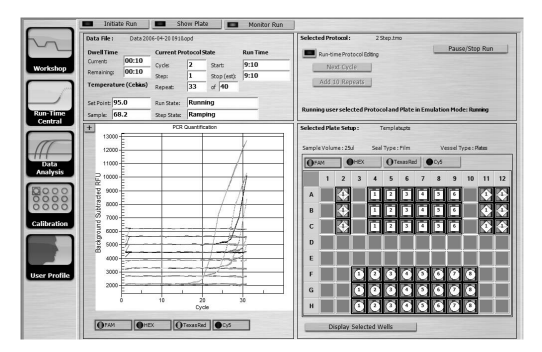
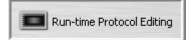


Fig. 4.6. The Monitor Run window.

4.4 Run-Time Protocol Editing

There are two choices available for modifying a protocol during a run: Next Cycle and Add 10 Repeats.

During a run, you can access these options by selecting the Run-time Protocol Editing check box as shown below.



Selecting this check box activates the Next Cycle and Add 10 Repeats buttons as shown below.



4.4.1 Next Cycle

Click Next Cycle to complete the current repeat of the present cycle before skipping to the next cycle. For example, you could use this feature when your samples have clearly crossed threshold and you want to skip to the melt cycle of your protocol.

4.4.2 Add 10 Repeats

Click Add 10 Repeats to add additional repeats to the current cycle. You can click this button multiple times, however, the total number of repeats is limited to 600. For example, it may be necessary to add repeats to a run in an experiment amplifying low copies of DNA to allow all samples to cross threshold. Click Add 10 Repeats to an amplification cycle of 30 repeats to make it 40 repeats.

Note: Modifications to the protocol are updated on the protocol displayed on the iCycler base module.

4.4.3 Pause/Stop Run

The Pause/Stop Run button allows you to pause a thermal cycling protocol. If you click Pause/Stop Run when the iCycler is at a setpoint temperature, the iCycler will hold at the setpoint temperature and stop counting down the dwell time. If you click Pause/Stop Run when the iCycler is ramping to the temperature, the iCycler will continue ramping until it reaches the next setpoint temperature, then pause at that step.

Clicking Pause/Stop Run activates two new buttons. Click Resume Run to resume the thermal cycling protocol. Click End Run to terminate the experiment.

Section 5 Data Analysis Module

The Data Analysis module is where data is presented and analyzed. When the iQ5 software opens, the Data Analysis module is gray and the module is not active. To analyze a data file, open the data file in the Data File tab of the Workshop module by selecting the data file and then clicking Analyze.

The Data Analysis module consists of six tabs:

PCR Quant

The PCR Quantification tab is used to set the analysis conditions for the data file. The analysis conditions include setting the PCR baseline, the threshold and which wells to exclude or include in the experiment. The analysis conditions should be set before using the Gene Expression, End Point or Allelic Discrimination tabs. For experiments with standards of known quantities, the PCR Quant tab is also where absolute quantities can be determined. The efficiency of the PCR reaction can also be determined using standard curves with either known quantities used to produce the standard curve or by using a serial dilution of the template under investigation.

Melt Curve/Peak

Melt Curve/Peak is a dynamic tool used to measure the melting temperature (T_m) of double stranded DNA molecules. DNA duplexes can be visualized by either incorporation of DNA-binding dyes (for example, SYBR Green I) or by hybridization with fluorescently labeled probes. Three major applications for Melt Curve/Peak are peak identification (number of amplified products), characterization of molecular beacons, and allelic discrimination.

End Point

End Point Analysis provides a convenient method of analyzing final RFU (Relative Fluorescence Unit) values. This can be useful when PCR Analysis is to determine if a given sample is Positive or Negative for a particular nucleic acid sequence.

Allelic Disc

The Allelic Discrimination tab is useful for assigning genotypes to unknown samples by making comparisons to known genotypes. It can be used to distinguish among homozygous wild types, homozygous mutants and heterozygous individuals based either on threshold cycle or on RFU.

• Gene Expr

The Gene Expression screen has flexible tools for the determination of the fold induction of one gene relative to another gene or relative to itself under different circumstances, e.g., temporally, geographically or developmentally different points.

• Edit Plate

In the Edit Plate screen, you may make changes to the sample type assignment or to the quantities of the Standards in the plate setup used to run the experiment. This is a simple way to salvage your experiment should you make mistakes in the sample type assignment. While you may make changes to the plate setup is never discarded and always remains with the file so that you may revert to it at any time.

5.1 PCR Quantification Tab

The PCR Quant tab is the tab that is first displayed after opening a data file that contains amplification data, as shown below. For experiments that lack amplification data, such as Melt Curve only or End Point Only experiments, the PCR Quant tab is grayed out and unavailable for analysis.

Use the PCR Quant tab to set the analysis conditions for the data file. The analysis conditions include setting the PCR baseline, setting the threshold, and determining which wells to exclude or include in the experiment. The analysis conditions should be set before using the Gene Expression, End Point or Allelic Discrimination tabs.

For experiments with standards of known quantities, the PCR Quant tab is also where you can determine absolute quantities for unknown samples. You can also determine the efficiency of the PCR reaction using standard curves with either known quantities used to produce the standard curve, or by using a serial dilution of the template under Investigation.

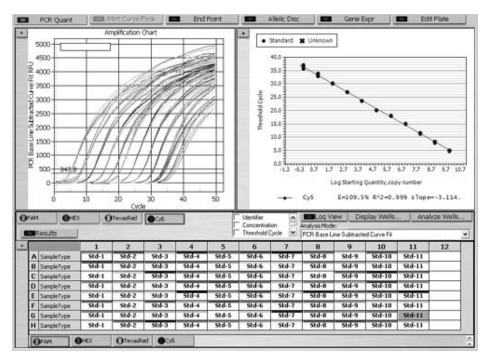


Fig. 5.1. The PCR Quant tab.

The PCR Quant tab consists of three panes:

- 1. Amplification Chart
- 2. Standard Curve Chart and
- 3. Results Section

5.1.1 Customizing the PCR Quant Display

You can customize the size of the panes in the PCR Quant tab in a number of ways. In the upper corner of each pane is a + (plus) button that enlarges the pane when clicked. The enlarged pane has a - (minus) button that reduces the pane when clicked. The amplification chart appears in the enlarged pane and includes an inactive - (minus) button when no standard curve is present. You can also move divider bars between each pane by clicking and dragging on the divider bar you want to move.

5.2 Amplification Chart

The Amplification Chart displays the relative fluorescence for each well at every cycle. Each trace represents the fluorescence of a given fluorophore for a single well and at each cycle a single data point is plotted which is the calculated mean of the data collected for that well during the particular cycle. The data that is used to determine this mean point is set by the Set Data Analysis Window dialog box. The data can be plotted in Background Subtracted, PCR Base Line Subtracted or PCR Base Line Subtracted Curve Fit mode.

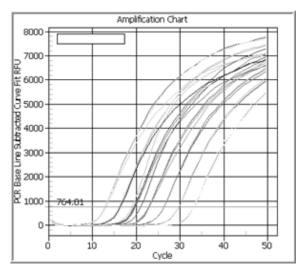


Fig. 5.2. The Amplification Chart.

5.2.1 Fluorophore Selector Buttons

Selecting Fluorophores to be Displayed

You can use the Amp Chart Fluorophore Selector buttons, which are located under the amplification chart, to display which fluorophores appear in both the amplification chart and standard curve chart. Selecting a single fluorophore to display is useful for determining the analysis parameters for that fluorophore. Selecting all fluorophores can be useful for ensuring that the efficiencies of each fluorophore set are approximately equal.

Use the fluorophore selector buttons to select and deselect which fluorophores to display. De-selecting a fluorophore does not remove the fluorophore from the analysis-only from the display.



Fig. 5.3. Fluorophore selector buttons.

Selected flurophore buttons display with a red border around the button perimeter. De-selected fluorophores display without this red border. In the example below, the FAM, HEX, and TexasRed flourophores have been selected, and Cy5 has been de-selected.

5.2.2 Select Analysis Mode

You can select from three analysis modes in the Analysis Mode drop-down list box as shown below.

Analysis Mode:	
PCR Base Line Subtracted	•

Fig. 5.4. The Analysis Mode drop-down list box.

The three analysis modes include:

- **Background Subtracted**: The background subtracted data is the relative fluorescence of each fluorophore after normalizing for exposure time, background factors, and well factors. No further analysis is possible on background subtracted data, therefore, the End Point, Allelic Disc, and Gene Expr tabs are unavailable.
- PCR Base Line Subtracted: To determine threshold cycles, construct standard curves, and determine the concentration of unknown samples, the data must be PCR baseline subtracted. The iQ5 software determines each PCR baseline subtracted trace by fitting the best straight line through the recorded fluorescence of each well during the baseline cycles. The iQ5 software then subtracts the best fit data from the background subtracted data at each cycle to generate the PCR baseline subtracted trace. By default, the software automatically chooses the beginning and end baseline cycles. You can override this default and manually give each trace a beginning and ending baseline cycle. User specified settings for PCR baseline subtraction may also be specified in the User Preferences module.

 PCR Base Line Subtracted Curve Fit: The iQ5 software fits the PCR baseline subtracted data to a smoothed curve using a balanced flank, centroid-finding digital filter. The curve fit process is performed in such a way that threshold crossing (C_T) is left invariant for all traces.

5.2.3 Log View Button

You can click Log View, shown below, to change between a semi-logarithmic and linear display of the Amplification Chart data.

The Log View button.



5.2.4 Setting Analysis Conditions for Threshold Cycle Calculation

In the amplification chart, the iQ5 software uses the two major analysis conditions that determine the threshold cycle as the baselines used for each individual trace and the threshold that is set for the fluorophore. By default, the iQ5 software calculates the baseline cycles and threshold when in PCR Base Line Subtracted or PCR Baseline Subtracted Curve Fit mode.

You can override the automatic conditions for baseline and threshold in the Base Line Threshold dialog box. Before you access this dialog box, ensure that only one fluorophore from the Amp Chart Fluorophore Selector buttons is selected, then right click on the Amplification Chart, and then click BaseLine Threshold. For other analysis condition settings, see the Amplification Plot Context Menu.

Analyze Wells

Click Analyze Wells, shown below, to select the wells that you wish to include in data analysis.

you click Analyze Wells, the Select Wells to Analyze dialog box appears as shown below.

🖉 Se	lect \	Vells	s to A	naly	ze			<u>.</u>				×
	AM		●н	EX		O Te	xasRe	d	Cv5	;		
	1	2	3	4	5	6	7	8	9	10	11	12
Α		2	J	•	J		,		J			12
В				1	2	3						
с				1	2	3						
D				1	2	3						
Е												
F			1	2	3	٩	5	6	0			
G			1	2	3	٩	5	6	0			
н			1	2	3	4	5	6	0			
	ted we					ellow	/black	c b ord	ers. Sel	ect/De:	select	
	Select	All		L	Ok		Ca	ancel		Appl	y	

Fig. 5.5. The Select Wells to Analyze dialog box.

You can include or exclude wells for analysis in this dialog box. The original data are always preserved and excluded wells can always be added back to the data analysis in this dialog box.

Removing wells from data analysis changes the calculation of the threshold location. Changing the calculation of the threshold location may result in a change in the Threshold crossing (C_{r}) location thus affecting all subsequent data analyses

that depend on C_{τ} values. This would include standard curve calculations and quantification of unknowns, Gene Expression, and Allelic Discrimination using Threshold Crossing values.

In addition removing wells from analysis will change the statistics for replicates.

To select the wells to include in data analysis:

- 1. In the PCR Quant tab, click Analyze Wells. The Select Wells to Analyze dialog box appears. A set of fluorophore selector |buttons appears at the top of this dialog box. The dialog box indicates the presence of a fluorophore in a well by that well having the fluorophore button color in the well. When you click a fluorophore button, the dialog box displays the sample type present in that well for that fluorophore.
- To select or deselect an individual well, click inside that well. Wells included in the analysis appear with a yellow and black border. Wells excluded from the analysis appear with a pale border (columns 10–12 in the above example). When a well is excluded from analysis, all fluorophores in that well are excluded.
- 3. To select all wells, click Select All.
- 4. To toggle the current selection so all wells currently selected will be unselected and vice versa, click the upper most left cell of the spreadsheet; this cell has the color of the currently selected fluorophore. You can also perform this action on a subset of the spreadsheet by clicking the appropriate row or column header. Only the wells in that row or column are toggled.
- 5. After you determine which wells to include in analysis, click Apply if you want the Select Wells to Analyze dialog box to remain open after the iQ5 software re-analyzes the wells. If you are satisfied with your selection, click OK to close the dialog box and update the wells to include in the analysis. If you click Cancel, the iQ5 software closes the dialog box and discards all changes.

Note: This procedure does not permanently remove data. This procedure only removes that data from the current analysis. You can add the excluded wells back to the analysis at any time by including them in the Select Wells to Analyze dialog box.

Display Wells

Click **Display Wells** to select the wells that you wish to include in the data display as shown below.

Displa	iy Wel	ls
--------	--------	----

After you click Display Wells, the Select Analyzed Wells for Display dialog box appears as shown below.

OF	AM		OH	x	_	OTes	casRec		DCv5	22	_	
	1	2	3	4	5	6	7	8	9	10	11	12
A												
B				1	2	3						
С				1	2	Þ						
D				1	2	J						
E												
F				0	٢	0	0	0	0			
6			0	2	٢	0	3					
н			0	2	0	0	3	0	0			
odifi		ils in a	highli II dye			illow/bl		rders.	Selec	t/Des		

Fig. 5.6. The Select Analyzed Wells for Display window.

You can include or exclude wells for display in the Select Analyzed Wells for Display dialog box. You can only choose from analyzed wells. Wells that have been excluded from analysis in the Select Wells to Analyze dialog box do not appear in the Select Analyzed Wells for Display dialog box.

Selecting wells for display does not change the underlying data analysis. Therefore, the calculation of thresholds and replicate statistics does not change.

To select the wells to include in data display:

- 1. In the PCR Quant tab, click Display Wells. The Select Analyzed Wells for Display dialog box appears. Only those wells that are included in the analysis appear in the dialog box. A set of fluorophore selector buttons appears at the top of this dialog box. The dialog box indicates the presence of a fluorophore in an analyzed well by that well having the fluorophore button color in the well. After you click a fluorophore button, the dialog box displays the sample type present in that well for that fluorophore.
- 2. Select or deselect an individual well by clicking inside that well. Wells included in the data display appear with a yellow and black border. Wells excluded from the data display appear with a pale border (rows G and H in the above example). When a well is excluded from analysis, the dialog box does not display the well (wells G8 and G9 in the above example).
- 3. To select all wells, click Select All.
- 4. To toggle the current selection so all wells currently selected will be unselected and vice versa, click the upper most left cell of the spreadsheet; this cell has the color of the currently selected fluorophore. You can also perform this action on a subset of the spreadsheet by clicking the appropriate row or column header. Only the wells in that row or column are toggled.
- 5. After you determine which wells to include in analysis, click Apply if you want the Select Analyzed Wells for Display dialog box to remain open after the iQ5 software re-analyzes the wells. If you are satisfied with your selection, click OK to close the dialog box and update the wells to include in the analysis. If you click Cancel, the iQ5 software closes the dialog box and discards all changes.

The Select Analyzed Wells for Display dialog box displays traces for fluorophores with parameters per the selected Amp Chart Fluorophore Selector buttons. You can use this dialog box to select all wells for display. Restore all traces for display by right clicking on the Amplification Chart and clicking Show All Traces in the menu.

Identifying a Specific Amplification Trace

Identify a specific trace by moving the mouse pointer along the trace until the hand icon appears. The dialog box identifies the trace, by both well name and fluorophore, in the top left corner of the Amplification Chart.

Selecting a Specific Amplification Trace

Select a specific trace by moving the mouse pointer on the trace until the hand icon appears, and then double click. The dialog box displays the selected trace. If you click Amplification Chart Fluorophore Selector, traces from other fluorophores in the selected well also appear.

Zoom

- To zoom in on a section of the plot, click and drag with the mouse on the desired region.
- To zoom out, select the plot and then type R, or right click on the plot and then click Restore Graph in the menu.

5.3 Standard Curve Chart

The Standard Curve Chart appears when, for a given fluorophore, more than 2 standards (with different quantities) are defined in the plate setup. The Amp Chart Fluorophore Selector buttons determine which fluorophores appear in the Standard Curve Chart, shown below.

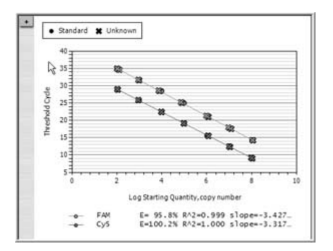


Fig. 5.7. The Standard Curve Chart.

The bottom of the chart displays a legend that includes:

- The color used to plot each fluorophore
- The name of the fluorophore
- The efficiency of the reaction
- The coefficient of determination (R^2)
- The slope of the line
- The y-intercept

You can enlarge the Standard Curve Chart by clicking on the plus (+) sign in the upper left corner of the pane.

5.3.1 Standard Curve Chart Menu

Right clicking on the Standard Curve Chart opens the Standard Curve Chart menu as shown below.

Copy Graph
Print Data
Print Graph
Restore Graph
Show Labels

Fig. 5.8. The Standard Curve Chart menu.

This menu includes the following options:

Copy Graph:	Copies the Standard Curve Chart to the clipboard. To copy the entire graph:				
	1.	Enlarge the chart by clicking the + button in the upper left corner of the Standard Curve Chart.			
	2.	Click Copy Graph in the Standard Curve Chart menu.			
Print Data:	Prints t Windov	he Standard Curve/Ct Results spreadsheet to your specified printer in vs.			
Print Graph:	Prints t	he graph to your specified printer in Windows.			
Restore Graph:		only after you zoom in on the chart. Clicking Restore Graph returns the rd Curve Chart to its un-zoomed state.			
Show Labels:	Labels	standards and unknowns with the well name as shown below.			

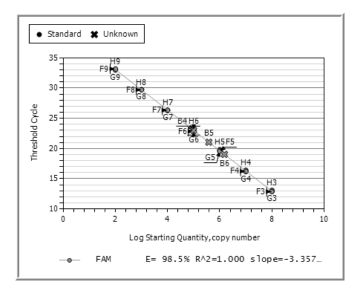


Fig. 5.9. The Standard Curve Chart with labels.

5.4 Results Section

The Results Workbook consists of three spreadsheets:

- 1. Plate Spreadsheet
- 2. Amplification Data (RFU) Spreadsheet and the
- 3. Standard Curve/Ct Results Spreadsheet

Click Results to toggle between the Plate Spreadsheet and the Amplification Data (RFU)/ Standard Curve/Ct Results spreadsheets.

Results

5.4.1 Plate Spreadsheet

The Plate spreadsheet displays data for each well in a grid fashion that represents the plate setup used in the experiment.

FAM	Results	ex [O TAMRA	TexasR	id Ocys		Concentration Threshold (End Point (lycle	Log Vi Analysis Mode PCR Base L		play Wells. d Curve Fit	Ana	lyze Wells.
		1	2	3	4	5	6	7	8	9	10	11	12
A	SampleType			Std-1	Std-2	Std-3	Std-4	Std-5	Std-6	Std-7	Std-8	NTC-1	
в	SampleType			Std-1	Std-2	Std-3	Std-4	Std-5	Std-6	Std-7	Std-8	NTC-1	
С	SampleType			Std-1	Std-2	Std-3	Std-4	Std-5	Std-6	Std-7	Std-8	NTC-1	
D	SampleType			Std-1	Std-2	Std-3	Std-4	Std-5	Std-6	Std-7	Std-8	NTC-1	
Ε	SampleType			Unkn-1	Unkn-2	Unkn-3	Unkn-4	Unkn-5	Unkn-6	Unkn-7	Unkn-8	NTC-1	
F	SampleType			Unkn-1	Unkn-2	Unkn-3	Unkn-4	Unkn-5	Unkn-6	Unkn-7	Unkn-8	NTC-1	
6	SampleType			Unkn-1	Unkn-2	Unkn-3	Unkn-4	Unkn-5	Unkn-6	Unkn-7	Unkn-8	NTC-1	
н	SampleType			Unkn-1	Unkn-2	Unkn-3	Unkn-4	Unkn-5	Unkn-6	Unkn-7	Unkn-8	NTC-1	

Fig. 5.10. The Plate spreadsheet.

The spreadsheet displays only one fluorophore at a time. You can use the Plate Spreadsheet Fluorophore Selector buttons found under the plate spreadsheet to determine which fluorophore displays in the spreadsheet. Each well has a colored bar at the top of the cell, which is the color used in the Amplification Chart to display the amplification trace for this well and fluorophore. The sample type is always displayed in this spreadsheet.

Above the spreadsheet is a set of check boxes you can use to display the Identifier, Concentration, Threshold Cycle, and End Point Call in the plate spreadsheet.

5.4.2 Amplification Data (RFU) Spreadsheet

You can view the individual RFU readings for each amplification trace at every cycle in the Amplification Data (RFU) spreadsheet. The Amp Data Fluorophore Selector buttons appear below the spreadsheet; these buttons control which fluorophore displays in the Amplification Data (RFU) spreadsheet. The spreadsheet can display data for only one fluor.

The well number appears at the top of the spreadsheet, and the cycle number appears down the side of the spreadsheet. In Single Point mode, each cycle is represented by one point that is the mean of all the data points analyzed during that cycle. In All Candidates mode, each individual data point displays.

5.4.3 Standard Curve/C₇ Results Spreadsheet

The Standard Curve/ C_{τ} Results spreadsheet displays the well, fluorophore, sample type, identifier, replicate number, threshold cycle, starting quantity and statistics for replicate groups. None of the data in this spreadsheet may be edited. The data can be copied to the clipboard for import into other software programs.

	Well	Fluor	Туре	Identifier	Replicate #	Threshold Cycle (Ct)	Log Starting Quantity	Starting Quantity (SQ)	SQ Mean	SQ Std. Dev.	Ct Mean	Ct Std. Dev	Set Point
69	H08	Cy5	Unkn		6	25.58	2.934	8.589e+002	8.66E+02	2.47E+01	25.57	0.041	N/A
70	H09	Cy5	Unkn		7	28.88	1.926	8.427e+001	9.20E+01	7.995+00	28.76	0.124	N/A
71	H10	Cy5	Unkn		8	31.36	1.165	1.466e+001	1.24E+01	1.72E+00	31.61	0.196	N/A
72	H11	Cy5	NTC		1	N/A	N/A	0.000e+000	0.00E+00	0.00E+00	0.00	N/A	N/A
73	A03	FAM	Std		1	12.17	8,000	1.000e+008	1.00E+08	0.00E+00	12.07	0.076	N/A
74	A04	FAM	9td		2	15.37	7.000	1.005e+007	1.00E+07	0.00E+00	15.30	0.125	N/A
75	A05	FAH	Std		3	18.65	6.000	1.000e+006	1.00E+06	0.00E+00	18.77	0.116	N/A
14	< > H	Amplificati	on Data (RFU), Star	dard Curve/ Ct Resu	ts/			•					

Fig. 5.11. Standard Curve Results Spreadsheet.

5.5 Amplification Plot Context Menu

You can access PCR amplification plot data analysis and display parameters by right clicking on the PCR amplification plot. After you right click on the plot, the amplification plot menu displays as shown below.

Set Data Analysis Window Digital Filter BaseLine Threshold
✓ Single Point All Candidates
Adjust Graph Define Trace Style Display Data Restore Graph Show All Traces
Copy Graph Print Graph Print Amplification Data Print Std Curve Data

Fig. 5.12. The amplification plot menu.

5.5.1 Data Analysis Options

There are three additional data analysis options in the amplification plot menu:

- Set Data Analysis Window
- Digital Filter
- BaseLine Threshold

Set Data Analysis Window

After you click Set Data Analysis Window... in the amplification plot menu, the Set Data Analysis Window appears as shown below. The number of data points collected per cycle depends on the exposure time and the dwell time of the cycle. You can change the percentage of data points used in determining the RFU value and you can choose to select this percentage from either the beginning or the end of the cycle or from a window in the middle of the cycle.

Set Data Analysis Window		\mathbf{X}
Locate Data Analysis Window		
Beginning of Cycle		
End of Cycle	Set Window Width (%	
Set Window Center and Width	Full cycle) :	99 🛨
Use Full Cycle Scan		
L	OK	Cancel

Fig. 5.13. The Set Data Analysis Window.

To set the Data Analysis Window:

- 1. Right click on the PCR amplification plot and then click Set Data Analysis Window in the amplification plot menu.
- 2. Click the Beginning of Cycle or End of Cycle option buttons to select data from the beginning of the cycle or the end of the cycle, respectively.
- 3. Set the window width in the Set Window Width scroll box.
- 4. To use all data points in the cycle select Use Full Cycle Scan.
- 5. Click OK to return to the PCR Quant screen.

Note: To center the analysis around data collected with a window in the middle of the cycle select the Set Window Center and Width radio button. Use the up or down arrows of the Set Window Width scroll box to select the percentage of data points. These data points will be chosen around the value set in the Window is Centered at % of Full Cycle text box. Click OK to return to the PCR Quant screen.

Digital Filter

After you click Digital Filter... in the amplification plot menu, the Set Digital Filters window appears as shown below. There are two intra-cycle data filtering options available: Rolling Boxcar and Weighted Mean. The default filter is the weighted mean as this filter is the only one

available during data acquisition.

Set Digital Filters	×
Intra Cycle	
Rolling Boxcar	
• Weighted Mean	
Disable All	
Enable Global Filter	Cancel

Fig. 5.14. The Set Digital Filters window.

The weighted mean is determined by the equation:

$Oi = (Ri + c^*M)/(1 + c)$

Where:

- Oi is the filtered value for a given data point, i
- Ri is the unfiltered value for data point i
- c is a weighted factor with a value of 2
- M is the arithmetic mean of all data points for the well within the given cycle.

The rolling box car filter is the arithmetic mean of the data readings i - (w-1), where w is the filter width.

The above filters only apply within a cycle. A global filter that smoothes data from cycle to cycle is also available by clicking the Enable Global Filter option button. The global filter operates on the trace for a given well and fluorophore using all cycles together in a single pass. Global filtering should be reserved for data that appears very noisy and should not be applied routinely.

BaseLine Threshold Settings

After you click BaseLine Threshold... in the amplification plot menu, the Base Line Threshold Parameter window appears and displays the fluorophore of the traces to be adjusted as shown below. In the example below, the window shows the FAM traces are to be adjusted.

		iold Paramete	r-FAM		
-	Line Cycle				
100	ito Calculate	0			
OUs	er Defined				
	Well	Fluor	Start	End	
1	AI	FAM	2	3	
2	A2	FAM	2	6	
3	A3	FAM	2	9	
4	A4	FAM	2	13	
5	A5	FAM	2	16	
6	A6	FAM	2	18	
7	A7	FAM	2	22	-
	10		-	~~	<u> </u>
	Sel	ect Ali	Edit Ra	ange	
* Ind	dicates chan	ged value			
Cros	sing Thres	hold			
0	Auto Calculat	ed Th	reshold Positio	in:	
•	Jser Defined	1	4.39		
			OK	Can	cel
			OK	Can	cel

Fig. 5.15. The BaseLine Threshold Settings Editor.

Manual Baseline Definition

Baselines are auto-calculated by default. You can override auto-calculation by clicking User Defined in the Base Lines Cycles area within the Base Line Threshold Parameter window. After you click User Defined, the iQ5 software activates the Select All and Edit Range buttons.

To edit a single trace, select the cell to the left of the well name, and then click Edit Range. Use the Edit Baseline Cycle Range dialog box to determine the beginning and ending baseline cycles. The dialog box indicates wells with manually determined baseline cycles with an asterisk.

Edit Baseline Cycle Range

The Edit Base Line Cycle Range dialog box has three modes.

- Both Start and Ending Cycle
- Start Cycle Only
- Ending Cycle Only

Both Start and Ending Cycle

In the Both start and ending cycle mode, you can edit both the Start Cycle and Ending Cycle boxes are editable. Enter the desired values for the start cycle and ending cycle for the selected traces and then click OK, as shown below.

dit		
Both start and ending o	tycle	•
Start Cycle	3	-
Ending Cycle	5	÷

Fig. 5.16. The Both start and ending cycle mode.

Start Cycle Only

In the Start cycle only mode, you can only edit the Start Cycle box. This mode is useful if you want to retain the automatically determined end cycle but wish to change the value of the start cycle. Enter the desired value for the start cycle for the selected traces and then click OK, as shown below.

dit		
Start cycle only	•	
Start Cycle	3 +	
Ending Cycle	5 🔅	

Fig. 5.17. The Start cycle only mode.

Ending Cycle Only

In the Ending cycle only mode, you can only edit the Ending Cycle box. This mode is useful if you want to retain the automatically determined start cycle but wish to change the value of the end cycle. Enter the desired value for the end cycle for the selected traces and then click OK, as shown below.

Edit		
Ending cycle only	•	
Start Cycle	3 -	
Ending Cycle	5 ÷	
	ОК Са	ncel

Fig. 5.18 The Ending cycle only mode.

Manual Threshold Definition

By default, the iQ5 software automatically calculates the threshold. You can override auto-calculation in one of two ways.

- 1. On the Amplification Chart, move the cursor over the displayed threshold line until the cursor icon becomes a hand. Left-click on the displayed threshold line and drag the threshold to the desired position.
- 2. From the Base Line Threshold Parameter dialog box, select the User Defined option in the Crossing Threshold section at the bottom of the dialog box. Enter the desired value for the threshold position and then click OK.

5.5.2 Data Display Options

There are seven additional data analysis options in the amplification plot menu:

- Single Point
- All Candidates
- Adjust Graph
- Define Trace Style
- Display Data
- Restore Graph
- Show All Traces

Single Point

The Single Point mode of data presentation, in which the iQ5 software averages all data collected at a particular step and then plots the average, is the default mode. For example, if the iQ5 software collects four data points during the third repeat of an amplification cycle, the mean of those 4 points is plotted at cycle 3. The alternative to viewing the data in Single Point mode is to view the data in All Candidates mode.

All Candidates

In this mode, the iQ5 software plots every single data point that is collected. You cannot use automated data analysis features when the data is in All Candidates mode. This mode is unavailable when the analysis mode is PCR Base Line Subtracted Curve Fit. The alternative to viewing the data in All Candidates mode is to view the data in Single Point mode.

Adjust Graph

You can re-scale or change the amplification plot from linear to log, or vice versa, by clicking Adjust Graph... in the menu. After you click Adjust Graph..., the Chart Axes Range Definition dialog box appears as shown below.

Chart Axes Range Definition	×
Y Max 5205 📑	
Y Min	X Max
0 📩	52 🛨
Restore Auto Scaled Chart	OK Cancel

Fig. 5.19. The Chart Axes Range Definition dialog box.

You can type the maximum and minimum values for either axis into the Y Max, Y Min, X Min, and X Max scroll box(es), or use the up and down arrows in each scroll box.

You can change the display to a semi-logarithmic view by selecting the Log View check box. Clear this check box to revert to a linear plot.

Revert the plot settings to the default by clicking Restore Auto Scaled Chart.

Note: You can also change the display to a semi-logarithmic view by selecting the Log View button in the PCR Quant tab.

Define Trace Style

You can customize the display by clicking Define Trace Style... in the menu. After you click Define Trace Style... the Define Trace Style dialog box appears as shown below.

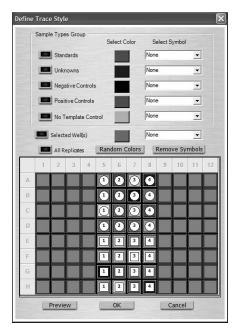


Fig. 5.20. The Define Trace Style dialog box.

You can change the trace color and symbol, which is used to display data points, on a well by well basis or in groups of sample types. You can also preview all changes before you apply them.

To modify a trace style:

- 1. Select the Sample Types Group to be modified, for example, all Standards. (Note that green is the default color that is automatically assigned to the Standards group when this button is active.)
- 2. To specify a different color for a given sample type group on the plate, click on the colored button in the Select Color column.
- 3. Select the symbol type (the default style is "None").
- 4. Click Preview to see your changes, and if those changes are satisfactory, click OK.

If you want to choose on a well by well basis, click the Selected Well(s) button, select a color and symbol type, and then select the desired wells in the plate grid at the bottom of the Define Trace Style dialog box. Click a column or row header to change an entire row or column. The dialog box outlines the edge of each well in the grid in the selected trace color. Select the All Replicates box to apply the selected color to all selected replicates of the wells.

To reset original trace style settings, click Random Colors and Remove Symbols, then click OK.

Display Data

Click Display Data to change the view of the results spreadsheet as follows:

- When Display Data is unselected, the results spreadsheet displays the plate spreadsheet view.
- When Display Data is selected, the results spreadsheet displays two tabs: the Amplification Data (RFU) tab and Standard Curve/Ct Results tab. You can perform the same function by clicking Results.

Restore Graph

Click Restore Graph to redraw the graph to its original size after you zoom in on the graph.

Show All Traces

Click Show All Traces to show all traces after you single out one or more using Display Wells or by clicking on a trace in the Amplification Plot.

5.5.3 Data Export Options

The Amplification plot menu offers four additional options for printing and exporting graphs or data:

- Copy Graph
- Print Graph
- Print Amplification Data
- Print Std Curve Data

Copy Graph

Click Copy Graph to copy the Amplification Plot into the clipboard to import into other programs.

Print Graph

Click Print Graph to print the graph to your default Windows printer.

Print Amplification Data

The Print Amplification Data option is only available when the iQ5 software displays the Amplification Data (RFU) spreadsheet. You can display this spreadsheet by clicking Results in the PCR Quant tab, or by clicking Display Data in the Amplification Chart menu.

When you click Print Amplification Data, the Print Preview dialog box appears so you can print the amplification data.

Print Std Curve Data

The Print Std Curve Data option is only available when the iQ5 software displays the Standard Curve/ Ct Results spreadsheet. You can display this spreadsheet by clicking Results in the PCR Quant tab, or by clicking Display Data in the Amplification Chart menu.

When you click Print Std Curve Data, the Print Preview dialog box appears so you can print the amplification data.

Note: If you alter the width of the columns in the spreadsheet in the Standard Curve/ Ct Results spreadsheet, the printed spreadsheet will reflect that change.

5.5.4 Exporting Results to Microsoft Excel

Export Data from the Results Table

- 1. Make sure the Results button is activated. If not, click Results.
- 2. Click the right mouse button. A shortcut menu appears. See the screen example below.
- 3. Select Export to Excel.
- 4. Enter a name and file destination for the Excel file generated with your data export.

	F	Results					
+		Well	Fluor	Туре	Identifier	Replicate #	Threshold Cycle (Ct)
-	1	B04	Cy5	Unkn	0 Hrs	1	18.22
	2	B05	Cy5	Unkn	1 Hr	2	18.09
1.000	3	B06	Cy5	Unkn	2 Hrs	3	18.05
	4	C04	Cy5	Unkn	0 Hrs	1	18.18
	5	C05	Cy5	UN .	rint	2	18.17
	6	C06	Cy5	Un	xport to Excel	3	18.03
-	7	D04	Cy5	Unkn	0 Hrs	1	18.23
	8	D05	Cy5	Unkn	1 Hr	2	18.20
	9	D06	Cy5	Unkn	2 Hrs	3	18.07

Fig. 5.21. Exporting PCR Quant Data tables to Microsoft Excel.

The Print command, on the Results table menu, will print the displayed Spreadsheet (Amplification Data, or Standard Curve/ C_{τ} Results). When selected, a Print Preview box is opened which contains an illustration of the spreadsheet as it will appear once printed. Clicking the printer icon opens the Windows print dialog box. Click OK to complete the printing task.

The Export to Excel command on the Spreadsheet menu is useful for exporting exact values from the spreadsheet. When the Export to Excel command is selected from the menu, an Export to Excel file save box is displayed. Choose a location of where the Excel file is to be saved and click Save. The iQ5 software automatically exports the selected data into a protected workbook.

- " The protected workbook generated by the iQ5 software contains the text values of what is represented on the Spreadsheet. For example, check boxes from the software application are replaced by "True" or "False" text in Excel.
- " The numeric values contained in the protected workbook are exact values from the software application that include several non-significant figures beyond the decimal point. This is important to note when considering whether to transfer iQ5 spreadsheet data by a copy and paste command or the Export to Excel command. With the copy and paste command only the significant digits displayed in the iQ5 software interface are transferred to Excel

5.6 Melt Curve and Melt Peak Charts

The RFU data collected during the melt curve part of the experiment are plotted as a function of temperature, as shown below.

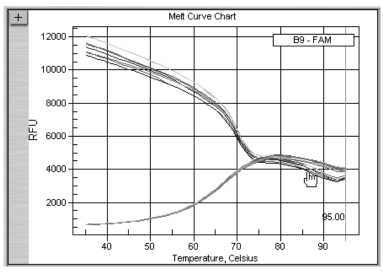


Fig. 5.22. Melt Curve Chart.

The vertical temperature bar may be dragged to any position on the plot. The temperature bar on the melt peak plot moves along to the same position as the temperature bar on the melt curve plot is moved. If you place the cursor over a trace, so that the pointer turns into a hand, the trace will be identified in the small box placed in the top right corner of the plot.

The chart can be expanded to fill the window by clicking the + box in the top left corner of the plot. Once expanded, it can be returned to its original size by clicking the - box in the top left corner.

Melt Peak Chart

The data in the melt peak chart are derived from the melt curve chart, as shown below.

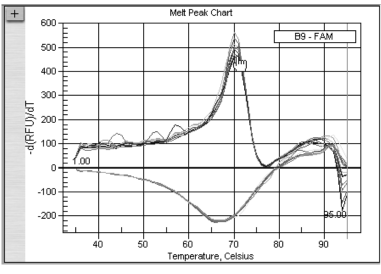


Fig. 5.23 Melt peak chart.

They represent the negative rate of change in fluorescence with changing temperature, that is:

-d(RFU)/dT

Where T is temperature.

In analyzing the melt peak data, the software assigns Begin and End temperatures to each peak and then calculates an area beneath that curve. The floor of the peak area is specified by the position of the threshold bar.

In order to be identified as a valid peak, a peak must have a minimum height relative to the distance between the threshold bar and the height of the highest peak. Therefore, if the threshold bar is dragged downward, increasing the

distance between the threshold bar and the highest peak, previously unidentified peaks may show up in the spreadsheet as their height becomes significant in relation to the highest peak. Similarly, dragging the threshold bar up can cause previously identified minor peaks to drop off the spreadsheet.

The vertical temperature bar may be dragged to any position on the plot. The temperature bar on the melt curve plot moves along to the same position as the temperature bar on the melt peak plot is moved. If you place the cursor over a trace, so that the pointer turns into a hand, the trace will be identified in the small box placed in the top right corner of the plot. The chart can be expanded to fill the window by clicking the + box in the top left corner of the plot. Once expanded, it can be returned to its original size by clicking the – box in the top left corner.

5.6.1 Melt Curve and Melt Peak Chart Menu

There are a number of features of the plot that may be modified as well as control of data filtering. All changes specified by the context menu accessed on the Melt Peak plot are also carried out on the Melt Curve plot. Those features are accessed by a right click on the chart, which brings up the following list:

Digital Filter				
Adjust Graph				
Define Trace Style Bostoro Graph				
Restore Graph Show All Traces				
Copy Graph				
Print Data				
Print Graph				

Fig. 5.24. Melt Curve/Peak Chart Context menu.

The menu options are:

- Digital Filter
- Adjust Graph
- Define Trace Style...
- Restore Graph
- Show All Traces
- Copy Graph
- Print Data
- Print Graph

5.6.2 Delete Selected Peaks

To remove a peak from the analysis, highlight the peak in the spreadsheet and then click Delete Selected Peaks as shown below. You can press the Shift key to highlight multiple peaks and delete them all at once.



5.6.3 Edit Melt Peak Begin/End Temperature

You may edit the begin and end temperatures for a melt peak by clicking the Edit melt peak begin/end temps option button as shown below.



After you click this option button, the currently-active peak displays alone with a Begin Temp bar and an End Temp bar. Drag the bars to the desired begin and end temperatures. As you drag the begin and end bars along the Melt Peak plot, iQ5 tracks the movement on the Melt Curve chart and updates the movement in the spreadsheet. When the Edit melt peak begin/end temps option button is activated, the Display Wells and Analyze Wells dialog boxes are unavailable.

In the figure below, the peak begin temperature is 55.5°C, the peak end temperature is 77°C, and the threshold is 1.

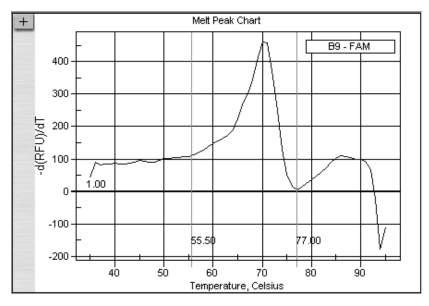


Fig. 5.25. Example Melt Peak Identification.

5.6.4 Melt Curve/Peak Control Area

The Control Area appears between the spreadsheet and the plots. This area contains buttons to access the Display Wells and Analyze Wells dialog boxes, as shown below.

	Display Wells		Analyze Wells		Restore Defaults
Temperature	Bar Thr	Threshold Bar		Selected We	ell Peak Height
55.5	0 -	- 1.0	00	B09	461.04

Fig. 5.26. The Control Area.

The position of the temperature bar, threshold bar, the height and well of the currently selected peak are also displayed here. You cannot edit any of these fields. When the Edit melt peak begin/end temp option button is active, the most-recently modified temperature (begin or end) displays in the Temperature Bar field.

The buttons at the top of the Control Area are:

- Display Wells
- Analyze Wells
- Restore Defaults

5.6.5 Melt Curve/Peak Spreadsheet

The spreadsheet below the plots displays information about the melt peaks, the RFU data or the -d(RFU)/dT data for each well. None of the fields may be edited, though you may eliminate peaks from the spreadsheet by either highlighting the peak in the spreadsheet and clicking Delete Selected Peak, or by dragging the threshold bar up to a position above the peak.

Identifier	Peak ID	Melt Temp.	Peak Desc.	Peak Height	Begin Temp	End Temp	Threshold Crossing Begin Temp	Threshold Crossing End Temp	Edited Begin Temp	Edited End Temp	
	B03.0	70.00		563.26	54.00	77.00	54.00	77.00	54.00	77.00	
	B04.0	70.00		538.08	54.00	77.00	54.00	77.00	54.00	77.00	
	B05.0	70.00		514.25	54.00	77.00	54.00	77.00	54.00	77.00	
	B06.0	70.00		490.74	60.00	77.00	60.00	77.00	60.00	77.00	
	B07.0	70.00		501.00	51.00	77.00	51.00	77.00	51.00	77.00	
	B08.0	70.00		469.23	55.00	77.00	55.00	77.00	55.00	77.00	
	B09.0	70.00		461.04	47.00	77.00	47.00	77.00	55.50	77.00	
	B10.0	71.00		504.03	38.00	77.00	38.00	77.00	38.00	77.00	
	Identifier	B03.0 B04.0 B05.0 B06.0 B07.0 B08.0 B09.0	Peak ID Temp. B03.0 70.00 B04.0 70.00 B05.0 70.00 B06.0 70.00 B07.0 70.00 B08.0 70.00 B08.0 70.00 B08.0 70.00 B08.0 70.00	Temp. Peak DD Temp. Peak Desc. B03.0 70.00 B04.0 70.00 B05.0 70.00 B06.0 70.00 B07.0 70.00 B08.0 70.00 B08.0 70.00	Temp. Peak Dest. Height B03.0 70.00 563.26 B04.0 70.00 538.08 B05.0 70.00 514.25 B06.0 70.00 490.74 B07.0 70.00 501.00 B08.0 70.00 469.23 B09.0 70.00 461.04	Deck DD Temp. Peak Dbsc. Height Temp. B03.0 70.00 563.26 54.00 B04.0 70.00 538.08 54.00 B05.0 70.00 514.25 54.00 B06.0 70.00 514.25 54.00 B06.0 70.00 501.00 51.00 B07.0 70.00 469.23 55.00 B08.0 70.00 461.04 47.00	Height Peak Des. Height Temp Find remp B03.0 70.00 563.26 54.00 77.00 B04.0 70.00 538.08 54.00 77.00 B05.0 70.00 514.25 54.00 77.00 B06.0 70.00 514.25 54.00 77.00 B06.0 70.00 510.00 51.00 77.00 B07.0 70.00 501.00 51.00 77.00 B08.0 70.00 649.23 55.00 77.00 B09.0 70.00 641.04 47.00 77.00	Identifier Peak ID Melt Temp Peak Desc. Peak Height Height Begin Temp End Temp Crossing Begin Temp 803.0 70.00 553.06 54.00 77.00 54.00 804.0 70.00 538.08 54.00 77.00 54.00 805.0 70.00 514.25 54.00 77.00 54.00 806.0 70.00 514.25 51.00 77.00 60.00 807.0 70.00 501.00 51.00 77.00 51.00 808.0 70.00 60.00 77.00 51.00 51.00 51.00 808.0 70.00 60.00 77.00 51.00 51.00 51.00 51.00 809.0 70.00 60.00 77.00 51.00 77.00 57.00	Identifier Peak ID Peak Term Peak Desc. Peak Height End Term Crossing Begin Crossing Begin Crossing End Term 803.0 70.00 532.65 54.00 77.00 54.00 77.00 804.0 70.00 538.08 54.00 77.00 54.00 77.00 805.0 70.00 54.00 77.00 54.00 77.00 54.00 77.00 806.0 70.00 54.00 77.00 54.00 77.00 54.00 77.00 807.0 70.00 54.00 77.00 54.00 77.00 54.00 77.00 808.0 70.00 51.00 51.00 77.00 51.00 77.00 809.0 70.00 64.00 77.00 51.00 77.00 77.00	Identifier Peak ID Metr Temp Peak Desc. Peak Height End Temp Temp Crossing Begin Temp Crossing Begin Temp Crossing End Temp End Temp Crossing Begin Temp Crossing End Temp End Temp Temp End Temp Crossing End Temp End Temp Temp Temp Temp Temp Temp Temp End Temp Temp	Identifier Peak ID Metrom Temp Peak Desc. Peak Height Temp Begin Temp Crossing Begin Temp C

Fig. 5.27. The Melt Curve spreadsheet.

The spreadsheet displays the following information for each peak:

Column	Description
Peak ID	A unique identification number in the format RCC.N where R is a row letter, CC is a column number, and N is a number beginning with 0, 1, 2, etc. In the example spreadsheet above, the first peak for well E3 is identified as E03.0 and the second peak as E03.1.
Melt Temp.	The temperature at the highest point of the melt peak.
Peak Height	The highest point of the melt peak.
Begin and End Temp.	Starting and ending point for melt peak. Area calculations are based on these starting and ending points. These values may be edited, but not in the spreadsheet.
Area	The area beneath the melt peak curve, bounded by the default peak begin and end temperatures and the default position of the threshold bar. Because this area calculation is defined by the default values of begin and end temperature and threshold bar, it does not change as the threshold bar is moved or if the begin or end temperatures are edited.
Area Fraction %	When there is more than one peak associated with a well, the contribution of each well to the total area beneath all melt peaks for that well is calculated. The iQ5 software makes this calculation using default begin and end temperatures and the default position of the threshold bar, so is not affected by edits to any of those parameters.
Edited Begin Temp.	When the peak begin temperature bar is dragged to a new position, it is reflected here.
Edited End Temp.	When the peak end temperature bar is dragged to a new position, it is reflected here.

5.7 Allelic Discrimination – For Multiplex Data Only

The Allelic Discrimination module is useful for assigning genotypes to unknown samples by making comparisons to known genotypes. It can be used to distinguish among homozygous wild types, homozygous mutants and heterozygous individuals based either on threshold cycle or on RFU. RFU data may be chosen from any cycle in the experiment. The assignments can be made automatically if controls are specified, or you can make the assignments manually.

5.7.1 Allelic Discrimination Plot

The allelic discrimination plot shows either RFU data or threshold cycle data from two different dye layers at the same time. Choose which plot to view using the RFU or Threshold Cycle Radio button in the Display Mode box.

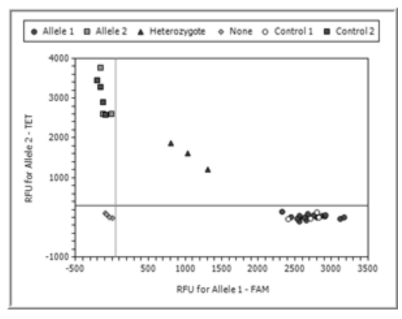


Fig. 5.28. Allelic Discrimination Plot, RFU view.

In Automatic Call mode, two bars, one vertical and one horizontal, divide the plot into four sections: one for each homozygous state, one for the heterozygous state and a non-reactive section. The positions of these bars may be adjusted in the Automatic Call mode. The bars do not appear in Manual Call mode.

Genotype assignments for unknown samples are determined by plotting the RFU for one fluorophore (allele 1 on the x-axis) against the RFU for the other fluorophore (allele 2 on the y-axis) on the allelic discrimination plot.

- If the unknown RFU values are greater than the horizontal bar and greater than the vertical bar, then the genotype is heterozygote.
- If the unknown RFU values are greater than the horizontal bar and less than the vertical bar, the genotype is homozygous for allele 2 (allele 2 RFU is plotted on the y-axis).
- If the unknown RFU values are less than the horizontal bar and greater than the vertical bar, the genotype is homozygous for allele 1 (allele 1 RFU is plotted on the x-axis).
- If the unknown RFU values are less than the horizontal bar and less than the vertical bar, then no genotype can be assigned.

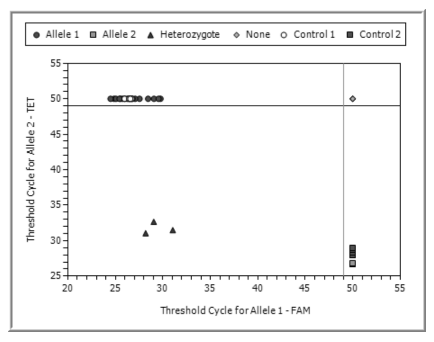


Fig. 5.29. Allelic Discrimination plot, Threshold Cycle view.

Genotype assignments for unknown samples may also be determined by plotting the threshold cycle for one fluorophore (allele 1 on the x-axis) against the threshold cycle for the other fluorophore (allele 2 on the y-axis) on the allelic discrimination plot.

- If the unknown threshold cycle values are greater than the horizontal bar and greater than the vertical bar, then the genotype is none. Samples not crossing threshold during the protocol are placed in this quadrant.
- If the unknown threshold cycle values are greater than the horizontal bar and less than the vertical bar, the genotype is homozygous for allele 1 (allele 1 threshold cycle is plotted on the x-axis).
- If the unknown threshold cycle values are less than the horizontal bar and greater than the vertical bar, the genotype is homozygous for allele 2 (allele 2 threshold cycle is plotted on the y-axis).
- If the unknown threshold cycle values are less than the horizontal bar and less than the vertical bar, then the genotype is heterozygote.

In Manual Call mode, the plot displays the RFU or threshold cycle data only. The threshold bars do not appear and modifications to calls are made manually by using the drop down menu in the Call column of the data spreadsheet or by choosing the call type from the radio buttons in the Allele Call box and then clicking and dragging to select desired samples in the graph.

You may zoom in on the allelic discrimination plot by first selecting Zoom Enabled from the Plot Menu, then left clicking and dragging over the area you wish to enlarge. Zoom out by choosing Restore Graph from the context menu.

5.7.2 Allelic Discrimination Plot Menu

Open the allelic discrimination plot menu by a right mouse click on the allelic discrimination plot as shown below.

Adjust Graph
Restore Graph
Zoom Enable
Show Labels
Copy Graph
Print Graph
Print Data

Fig. 5.30 The Allelic Discrimination plot menu.

- Adjust Graph. This feature allows you to change the way that the plot is presented. The maximum and minimum values for either axis may be entered directly into the text boxes. Alternatively the up and down arrows can be used to set the maximum and minimum values.
- Restore Graph. Use this to redraw the graph after zooming.
- Zoom Enable. Use this to enlarge the desired plot area.
- Show Labels. Labels each sample on the Allelic Discrimination Plot with its well name.
- Copy Graph. This will copy the displayed Allelic Discrimination plot to the clipboard for import into other programs.
- Print Graph. This prints only the graph.
- Print Data. This prints the Allelic Discrimination Spreadsheet data.

5.7.3 Allelic Data Spreadsheet

A six-column spreadsheet is displayed in the window as shown below. The first column is the well number, and the second and third are the Identifier entered at plate setup for the two dye layers. The fourth and fifth column present the RFU or Threshold Cycle data for both dye layers and the last column is the genotype call made by either the software or the user.

	ID 1	ID 2	RFU 1	RFU 2	Call	
B1	Control WT	Control WT	-1109.2	3305.54	Control2	
B2	Control WT	Control WT	-841.64	3443.69	Control2	
B3	Control WT	Control WT	-527.05	2935.91	Control2	
B4	Control WT	Control WT	-384.4	2604.26	Control2	
B5	Control WT	Control WT	2446.98	-97.95	Control1	
B6	Control WT	Control WT	2829.31	-103.97	Control1	
B7	Control WT	Control WT	2862.07	42.02	Control1	
B8	Control WT	Control WT	2767.72	3.35	Control1	
D1	NTC	NTC	-74.64	66.58	None	
D2	NTC	NTC	-67.28	-97.83	None	
D3	NTC	NTC	-64.89	-142.04	None	
D4	NTC	NTC	-151.4	-36.8	None	
E1			-1278.73	3751.75	Allele2	
E2			802.44	1860.93	Heterozygote	
53			1109.11	1573.08	Heterozvaote	–

Fig. 5.31. The allelic data spreadsheet.

- In Automatic Call mode, the iQ5 software creates the assignments based on the positions of the vertical and horizontal bars.
- In Manual Call mode, the Call column becomes editable through a menu. This can be very useful if you want to change the definition of a sample from an unknown to a positive control, for example.
- The data in the spreadsheet may be copied to the clipboard for export to another program by clicking in the top left corner of the spreadsheet and typing CTRL+C.

5.7.4 Automatic/Manual Call

Select the type of analysis by clicking one of the following option buttons.



Fig. 5.32. The Automatic Call and Manual options.

Automatic Call is the default.

- Choose Automatic Call for the software to make genotype assignments for every unknown based on the positions of the vertical and horizontal bars and presents those assignments in the data spreadsheet. If at least three positive (homozygous) controls are specified for each dye layer, the positions of the bars are based on the mean and standard deviation of the threshold cycles or RFU values of the controls. If insufficient numbers of controls are specified, then the position of each bar is determined by the range of threshold cycles or RFU values in the selected dye layers. The positions of the bars may be manually adjusted by dragging them. Click Recalculate after any change in the position of the vertical and horizontal bars and the software will make new genotype assignments based on the adjusted positions.
- Click Manual Call to make the genotype assignments directly into the data spreadsheet or on the plot. You can use the manual call feature to change the definitions of one or more wells on the plot or in the spreadsheet and then return to Automatic Call and the software will position the vertical and horizontal bars based on the new definitions, and then make genotype assignments.

5.7.5 Manual Calls

To make manual calls on the plot.

1. Click Manual Call. A new box appears in the window.

Control 1
Control 2
-
Unknown

Fig. 5.33. The Manual Call specification options box.

- 2. Select one of the Allele Call types by clicking its radio button.
- 3. On the plot, drag the cursor around the well or wells to be assigned the call. When the mouse button is released, the new assignment will be made on both the plot and the data spreadsheet.

To make manual calls in the data spreadsheet.

- 1. Click Manual Call.
- 2. Click the cell for the desired well in the Call column of the data spreadsheet. Select the new call from the pull down menu.
- 3. Note that "unknown" may only be selected in manual call mode.

5.7.6 Display Mode

Choose which aspect of the data will be analyzed by clicking one of the following option buttons.

Display Mode	
Threshold Cycle	
ORFU	

Fig. 5.34. The Display Mode option buttons.

When you click RFU, the select cycle option appears so you can analyze RFU data from any cycle. Change the cycle by entering a new value in the Select Cycle drop down list box as shown below.

Select Cycle:	40	-
---------------	----	---

Fig. 5.35. The Select Cycle drop down list box.

5.7.7 Vertical Threshold

The Vertical Threshold box, shown below, displays the current position of the vertical bar. This box is not editable. The positions of the vertical bar along with the position of the horizontal bar determine the genotype assignment for unknown samples.

Display Mod	e
Threshold	dCycle
ORFU	

Fig. 5.36. The Vertical Threshold box.

5.7.8 Horizontal Threshold

The Horizontal Threshold box, shown below, displays the current position of the horizontal bar. This box is not editable. The position of the horizontal bar along with the position of the vertical bar determines the genotype assignment of unknown samples.



Fig. 5.37. The Horizontal Threshold box.

5.7.9 Normalize Data

When the RFU option button is selected, the Normalize Data option button appears as shown below if the plate setup contains a no template control (NTC) sample type for both Allele 1 and Allele 2.



Fig. 5.38. The Normalize Data button.

The RFU data may be normalized and displayed on a plot that ranges from 0 to 1 on both axes. This display is sometimes a very effective presentation of RFU data. The RFU data is normalized to the NTC value as a linear combination of Allele 1 and Allele 2 specific signals. The iQ5 software uses the following formula:

Normalized
$$A_{I} = \frac{A_{I}}{A_{I} + A_{2} + \overline{\chi}(NTC_{AI} + A_{2})}$$

Where, A_1 represents Allele 1 A_2 represents Allele 2 \overline{x} represents the mean NTC_{A1+A2} represents the sum of NTC for Allele 1 and Allele 2 Reference:

Livak JL, Marmaro J and Todd JA, Towards fully automated genome-wide polymorphism screening, Nature Genetics, 9, 341–342 (1995)

5.7.10 Restore Default

At any time, all modifications to the allelic discrimination data can be reversed by clicking Restore Default. The iQ5 software will reload the original X- Axis allelic 1 fluorophore and Y -Axis allelic 2 fluorophore data, original Display mode and Show Label setting which were originally saved in OPD file and will also set Automatic Call Method to recalculate the data, plot the chart and display the results in the allelic discrimination data spreadsheet .

5.8 End Point Analysis

The End Point Analysis module is a convenient method of analyzing final RFU, or Relative Fluorescence Unit, values.

Endpoint analysis can be performed in two ways:

- Selecting the Endpoint tab in the Data Analysis module for an existing data file
- Click Run End Point to initiate the collection of end point data from a sample plate.

5.8.1 End Point Analysis of Existing Data Files

Within the Data Analysis module, any iQ5 or MyiQ data file with PCR quantification data can be viewed in the End Point tab. However, the corresponding PCR quantification data must be analyzed in either the PCR Base Line Subtracted Mode or the PCR Base Line Subtracted Curve Fit mode before the End Point tab becomes available. Furthermore, you may choose to include only certain wells under either tab, since well selection is synchronized between the PCR Quant tab and the End Point tab.

5.8.2 Initiating an End Point Run

When initiating an End Point run, two repeats of a thirty-second step will be run. Programmed to collect only end point analysis data, only the setpoint temperature of the End Point protocol can be modified. To initiate the collection of end point data from a sample plate:

- 1. Insert the plate into the iCycler reaction module.
- 2. Create or open a plate setup file.

Note that any positive or negative controls defined in the loaded plate setup will be exported to the endpoint analysis table after the run. You may always define all your controls post-run from the End Point or the Edit Plate tab in the Data Analysis module.

- 3. Click on Run End Point. The Initiate Run tab appears.
- 4. Specify the setpoint for data collection in the Setpoint box.

Note: You must use Persistent Well Factors for every End Point Analysis Run.

5. Click Begin Run, give the data file a name, and the end point analysis run will begin. Upon the completion of end point data collection, the End Point tab appears and displays the results.

5.8.3 Analyzing Endpoint Data

End point data is displayed immediately following an End Point run. Although any file with amplification data may be analyzed post-run in both the PCR Quantification and End Point Analysis tabs, an end point analysis run may only be analyzed in the End Point tab. Melt curve-only experiments, which neither contain amplification data nor end point data, may not be analyzed in the End Point tab.

The End Point tab is comprised of several sections; each one described in detail below.

End Point Analysis Methods

Below the file and fluorophore information is the Method box. The Method box allows you to select the method of assigning positive and negative values to your unknowns based on RFU values. The Method box consists of the following three choices:

- **Negatives**: This is the default method. Select this method to use negative controls to define/call unknown samples. Samples are considered positive if they are greater in RFU value than the negative control average plus the tolerance.
- **Positives**: Select this method to use positive controls to define/call unknown samples. Samples are considered positive if they are greater in RFU value than the positive control average minus the tolerance.

• **Positives & Negatives**: Select this method to use positive and negative controls to define/call unknown samples. Samples are considered positive if they are greater in RFU value than the positive control average minus the tolerance. Samples are considered negative if their RFU value is less than the negative control average plus the tolerance.

End Point Tolerance

End Point Tolerance defines the margins for sorting unknowns as positives or negatives. How the tolerance variable and the type of tolerance are applied depends on which Method is selected. The End Point Tolerance drop-down list box consists of two choices:

- **RFUs**: This is the default Tolerance choice and should be selected if you would like to use an absolute RFU value for the tolerance value. The minimum RFU tolerance value is 2, whereas the maximum is the absolute value of the highest RFU value minus the absolute value of the lowest RFU value. The default RFU tolerance value is 10% of the total RFU range.
- **Percent of Range**: Select this setting if you would like to use a percentage of the RFU range for the tolerance value. The minimum percent of range is 1 percent, whereas the maximum percent of range is 99 percent. The default percent of range tolerance is 10 percent.

The definition of the RFU range is dependent on the Method chosen.

- For the Negatives method, the range is the highest RFU value minus the Negative control average.
- For the Positives method, the range is the positive control average minus the Lowest RFU value.
- For the Positives & Negatives Method the range is the positive control average minus the Negative control average.

Below the Method box, are the End Cycles to Average and Number of Ranks boxes as well as the Sort Data by Call check box.

End Cycles to Average

End cycles to Average is the number of cycles from the last cycle, that will be used to calculate an average End Point RFU value. The end cycles to average field defaults to 2 for End Point Only runs, and 5 for non-End Point Only runs. In order for the iQ5 software to analyze non-End Point Only data in the End Point Analysis tab, at least 6 repeats at data acquisition must be performed.

Number of Ranks

The Number of Ranks allows assignment of samples into distinct groups based on their RFU values. The absolute range (Highest RFU minus Lowest RFU) is divided by the number of ranks selected. The default rank value is 10 and the minimum number of ranks is 3.

Colored rank boxes, displayed below Number of Ranks and Sort Data by Call, symbolize the number and order of ranks in the End Point Analysis. To the right of the colored rank boxes are five color-gradation buttons that allow a change in the color scheme of the rank boxes, once the data are analyzed.

Sort Data by Call

This function is used to sort End Point Samples into Positive and Negative Calls (as well as No Calls wells which do not fall into the previous two categories). Sort Data by Call is essentially the Positives & Negatives method but also includes sorting and color coding the Positive and Negative Calls. The ranking function is disabled in this mode.

Results

The Results box lists the following information:

- **Source of Data**: Displays the analysis mode of the source data. For End Point Only Data this is Background Subtracted. For Endpoint Analysis of PCR Quantification runs the source of data must be either PCR Base Line Subtracted or PCR Base Line Subtracted Curve Fit. You may alter the analysis mode of the source data in the PCR Quantification tab within the Data Analysis module.
- Current method: Displays the currently selected method for assigning positive and negative values to unknowns.
- **Tolerance**: Displays the currently selected mode of tolerance calculation. Immediately beneath the displayed mode is the tolerance value. If RFUs are chosen then the Set Tolerance in RFUs is given. If Percentage of Range is chosen then the Calculated Tolerance in RFUs appears.
- **Range**: The value displayed here is dependant on the Method chosen.
 - If the Positives method is chosen then the Range is the Positive Controls Average RFU minus the Lowest RFU.

- If the Negatives Method is chosen then the Range is the Highest RFU minus the Negative Controls Average RFU.
- If the Postives & Negatives method is chosen then the Range is the Postive Controls Average RFU minus the Negative Controls Average RFU.
- + Controls Average RFUs: Displays the average RFU of all positive controls. No value is shown if there are no
 positive controls defined.
- **+ Control Tolerance**: In the Postives or Positives & Negatives Methods, samples with values equal to or higher than this amount will be called as Positive. No value is shown if there are no positive controls defined.
- - Controls Average RFUs: Displays the average RFU of all negative controls. No value is shown if there are no negative controls defined.
- - Control +Tolerance: In the Negatives Method, samples with values equal to or higher than this amount will be called as Positive. In the Positives & Negatives Method, samples with values less than this amount will be called Negatives. No value is shown if there are no negative controls defined.

5.8.4 End Point Analysis Spreadsheet

The End Point Analysis Spreadsheet contains the necessary data to perform End Point analysis, and also displays the values assigned to unknowns after the Recalculate button is pressed. The table headings include the following:

- Well: Lists the well ID of each sample. You may click on the Well column header to sort the table by wells. Note that you may also include or exclude certain wells from End Point analysis via the Analyze Wells button above the spreadsheet.
- **Sample Type**: Lists the sample type of every well, as defined in the data's plate setup. You may click on the Sample Type column header to sort the table by sample type.
- End RFU's: Lists the absolute RFU averages for each well, as calculated from the end cycles, which are specified in the End Cycles to Average field. You may click on the End RFU's column header to sort the table by end RFUs.
- **Define Controls**: Lists any positive or negative controls defined in the original plate setup file. You may also add new controls or edit existing controls in this column by clicking on the drop down menu on the right side. The options are (+) Positive, (-) Negative, or blank. Alternatively, you may type the letter 'p' or the plus sign (+) to select a positive control, and you may also type the letter "n" or the minus sign (-) to select a negative control. The controls specified in this column are used in the End Point analysis calculations that assign positive or negative values to the unknowns. You may click on the Define Controls column header to sort the column by the controls.
- **Unknowns Call**: If a well is not defined as a positive control or a negative control in the Define Controls column, it is considered an unknown for End Point analysis. This column displays the value assigned to each unknown after the Recalculate button has been pressed. Unknowns Call may be (+) Positive, (-) Negative, or blank. You may click on the Unknowns Call column header to sort the table by Unknowns Call.
- **Unknowns Ranking**: Lists the rank into which each unknown falls. An unknowns ranking depends on the total number of ranks and the End RFU's value. You may click on the Unknowns Ranking column header to sort the table by Unknowns Ranking.
- **Identifier**: Lists the identifier for every well, as defined in the data file's plate setup. You may click on the Identifier column header to sort the table by identifier.

5.8.5 Recalculate Button

Clicking Recalculate performs the final calculations used to determine unknown sample calls. The Recalculate button is active when sufficient controls have been specified for the selected Method and End Point Tolerance settings. The following conditions result in an error message:

- If the Positives method is selected and there is not at least one positive control defined in the End Point analysis spreadsheet.
- If the Negatives method is selected and there is not at least one negative control defined in the End Point analysis spreadsheet.
- If the Positives & Negatives method is selected and there is not at least one positive control and at least one negative control defined in the End Point analysis spreadsheet.
- If no Controls are defined.
- If the range is less than one (that is, the negative control RFU average is larger than the positive control RFU average).

End Point analysis cannot be performed if one or more of these error conditions is present when you click Recalculate.

After you are satisfied with your End Point analysis, you may choose to view, save, or print a customized End Point Analysis report. Click the Reports menu to obtain customized reports for the End Point data. Click Print to print the End Point spreadsheet.

5.9 Gene Expression Analysis

The Gene Expression screen has flexible tools for the determination of the fold induction of one gene relative to another gene or relative to itself under different circumstances, i.e. temporally, geographically or developmentally different points. Additionally, within Gene Expression it is possible to create Gene Studies from multiple plates performed at different times.

Gene Expression Analysis Overview

Activate the Gene Expression tab from the Data Analysis Module by clicking Gene Expr as shown below.

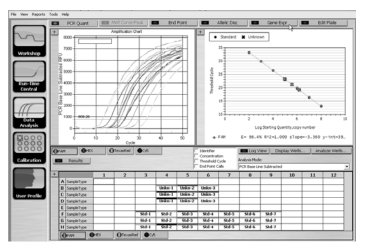


Fig. 5.39. Preparing Amplifcation Data for Gene Expression Analysis.

You can use the Gene Expression tab of the iQ5 software to evaluate relative differences in any target concentration. For example, you can evaluate relative numbers of viral genomes or stably transfected sequences. The most common application is evaluating target concentration in cDNA samples to infer steady state messenger RNA levels.

Gene expression analysis is the most common real-time PCR application. Therefore, this manual uses terminology that is specific to this kind of study. You can safely substitute "gene" with the word "sequence" or "target" when you read about how to use the Gene Expression module.

Without stringently quantified controls, you cannot use this feature of the iQ5 software to evaluate target concentrations of particular sequences relative to each other. The iQ5 software can only evaluate relative differences of a sequence between a group of samples.

Basic Workflow Steps

- 1. Open a Bio-Rad Real-Time PCR Data file.
- 2. Assess the Threshold and Baseline for the data file and then make changes if necessary.
- 3. Click the Gene Expression tab.
- 4. Make any changes to the Genes and Condition (for example, Sample and Treatment) assignments in the Gene Expression Plate interface.
- 5. Choose an analysis method:
 - a. Normalized Expression ($\Delta\Delta$ C_T) is the default; or
 - b. Relative Quantity (ΔC_T).
- 6. Assign attributes (sample color, show graph, etc.) in the Gene List.
- 7. Assign attributes (sample color, show graph, etc.) in the Condition List.
- 8. If you choose Normalized Expression ($\Delta\Delta$ C₇), assign reference gene(s).

9. Click Recalculate to see your results.

If you need to compare these data to results obtain in other .opd files, you will need to enable this file for Multi-file Gene Expression analysis, also called a Gene Study. To enable your file for Gene Study:

10. Click the Enable for Gene Study button.

11. Go to the File menu to save your file.

Normalized Expression is graphed in the Gene Expression Graph interface. The data can be accessed in the Gene Expression Settings interface within the Data Table.

5.9.1 Using Reference Genes (Normalized Gene Expression, $\Delta\Delta C_{T}$)

Rather than using some other method to normalize data, you may use one or more reference gene's message levels as assayed in your qPCR experiment as a normalization factor. Reference genes should be genes which are not regulated in the biological system being studied.

Normalized expression is the relative quantity of your gene normalized to the relative quantities of the reference gene(s). Simply stated, the quantities of the reference genes are used to normalize the values of your genes of interest.

Provided the reference genes are not regulated in your system this will account for loading differences or variations in cell # represented in each of your samples. This value is sometimes referred to as $\Delta\Delta C_T$ because of the equation initially introduced by Livak, et al., to evaluate normalized expression. The software uses a modification of this equation. View the equation used to calculate normalized expression.

Normalized Expression can be viewed as normalized target sequence quantity if you are using the software for other purposes.

Workflow for Normalized Gene Expression ($\Delta\Delta C_{\tau}$) Analysis

The workflow for normalized gene expression analysis is:

- 1. Open a Bio-Rad Real-Time PCR Data file.
- 2. Assess Threshold and Baseline for the data file and make changes if necessary.
- 3. Click the Gene Expression tab.
- 4. Make any changes to Genes and Condition (for example, Sample and Treatment) assignments in the Gene Expression Plate interface.
- 5. Make sure the Normalized Expression ($\Delta\Delta C_T$) button is activated. If not, click Normalized Expression ($\Delta\Delta C_T$).
- 6. Assign reference genes in the Gene List.
- 7. Click Recalculate to see your results.
- 8. To compare results to data from other assays in a Mutlifile Gene Expression study, click Enable for Gene Study, and save your file.

5.9.2 Not using Reference Genes (Relative Quantity, ΔC_{T})

By definition, Relative Quantity data is not normalized. Typically researchers that do not use reference genes are confident in one of the two following considerations.

- 1. Each condition (sample) assayed represents the same amount of biological sample. Typically they choose to load the same mass of RNA or cDNA in each well and feel that mass of nucleic acid is an effective way of normalizing the resulting data. No modification of Relative Quantity data is needed too obtain normalized data. The data is normalized by experimental design.
- Any variance in the amount of biological sample loaded will be normalized in post PCR analysis by some method. For example, a researcher might choose to simply divide the relative quantity value by the normalizing factor indicated after each of the examples listed below. Options may include but are not limited to:
 - a. Mass of nucleic acid loaded for each sample Rel Quant/ng RNA represented in each sample
 - b. Number of cells from which nucleic acid was isolated Rel Quant/number of cells represented in each sample
 - c. Mass of tissue from which nucleic acid was isolated Rel Quant/mass of tissue represented in each sample

Relative Quantity is the target sequence concentration relative to other samples in the experiment. This is sequence quantity without taking into account reference genes for normalization. No calculations are made to account for loading differences or differences in the number of cells represented in each of your samples. Relative Quantity can be viewed as Non-Normalized Expression. This value is sometimes referred to as ΔC_T because of the equation used to calculate relative quantity.

Workflow for Relative Quantity (ΔC_{T}) Analysis

- 1. Open a Bio-Rad Real-Time PCR Data file.
- 2. Assess Threshosld and Baseline information for the data file and make changes if necessary.
- 3. Click the Gene Expression tab.
- 4. Make any changes to Genes and Condition (for example, Sample and Treatment) assignments in the Gene Expression Plate interface.
- 5. Click Relative Quantity.
- 6. Click Recalculate to see your results.
- 7. To compare results to data from other assays in a Mutlifile Gene Expression study, click Enable for Gene Study, and save your file.

5.9.3 Specifying Gene and Condition Labels for Gene Expression Analysis

Open a Bio-Rad Real-Time PCR Data file.

- 1. Assess Threshold and Baseline for the data file and then make changes if necessary.
- 2. Click the Gene Expression tab.
- 3. Highlight the wells in the gene expression plate interface which you wish to edit.
- 4. Change the gene and condition assignments in these wells by typing in the gene pull-down menu and the condition pull-down menu. You may use the default values which already populate these lists if you like.

Assigning Gene and Condition Names

To access the gene list:

- 1. Make sure the Settings button is activated. If not, click Settings. See the screen example below.
- 2. Click the Gene List option button.

The Gene List appears in the spreadsheet directly below the Gene List option button as shown below.

All	elic Disc		Gene E	¢pr	
-	Setting	Data Tabl	e		
G	ene List	Condition List	• Data S	et List	
	Name	Full Name	Ref	Color	Show Graph
1	Tubulin	Tubulin			~
2	Actin	Actin	v		~
3	IL1b	IL1b			~
4	Gapdh	Gapdh	~		•

Fig. 5.40. Selecting the Gene List Option.

Assigning a Reference Gene

To assign a reference gene:

- 1. Click Gene List. See the screen example below.
- 2. Select the check box in the Ref column within the control condition(s) you want to assign a reference gene to. You can assign the role of reference gene to as many genes as you wish.

-	Setting	Data Tab	le				
Ge	ene List	Condition List	• Data S	et List			
	Name	Full Name	Ref	Color	Show Graph	Auto Efficiency	Efficiency (%)
1	Tubulin	Tubulin			v	v	95.8
2	Actin	Actin			~	v	95.9
3	IL1b	IL1b			~	V	94.6
4	Gapdh	Gapdh			~	V	94.7

Fig. 5.41. Assigning a Reference Gene.

Condition List Option

You can view conditions as samples in the Condition List spreadsheet as shown below. They are particular tests or conditions being evaluated for the purposes of your experiment.

A condition can be as simple as "sample 1" or complex as "mouse #123558 liver + PMA", though the latter example is often too long to present aesthetically on a graph.

Control Condition

You can assign one condition as the control condition by selecting a check box next to the sample in the Condition List.

The iQ5 software assigns the control sample a value of 1 for every gene and all other conditions will be presented with values relative to this one. This makes it simple to evaluate fold expression relative to the chosen (control) sample.

Note: The control condition will have a value of one for all genes.

Accessing the Condition List

To access the condition list:

- 1. Make sure the Settings button is activated. If not, click Settings. See the screen example below.
- 2. Click the Condition List option button.

-	Setting	Data Tab	le			
• Ge	ene List 🛛 🔘	Condition List	• Data S	et List		
	Name	Full Name	Ctrl	Color	Show Graph	
1	Std-3	Std-3				
2	0 Hrs	0 Hrs	R		v	
3	Std-5	Std-5	L/3			
4	Std-4	Std-4				
5	1 Hr	1 Hr			v	
6	Std-6	Std-6				
7	Std-7	Std-7				
8	2 Hrs	2 Hrs			V	
9	Std-1	Std-1				
10	Std-2	Std-2				

Fig. 5.42. Accessing the Condition List.

Assign a Control Condition

To assign a control condition:

- 1. Click the Condition List option button. See the screen example below.
- 2. Select the check box(es) in the Ctrl column next to the condition(s) which you want to assign as control conditions.

5.9.4 Data Set List

The Data Set List displays pertinent information about the data being analyzed in a given Gene Study or .opd file. The information contained within the cells of the Data Set List is for display only, and cannot be modified.

Each row of the Data Set List is linked to a single data tab of the Gene Expression Plate Interface, located at the bottom of the Gene Expr window. Selecting a row from the spreadsheet automatically toggles to its representative dye-layer data tab in the Plate Interface. The inverse is also true, that is, selecting a dye-layer data tab results in toggling to its representative row in the Data Set List spreadsheet.

Information displayed in the Data Set List includes:

- **Name**: Displays the fluorophore name for a given set of collected data. A number is placed before the abbreviated fluorphore name. The assigned number helps to distinguish between data from the same fluorophore when multiple files are used (as in a Gene Study).
- **Full Name**: This cell contains the original name of the opd file as well as the original location of the opd when it was created.
- Created Date: This cell displays the date and time the .opd file was created.

The Name and Full Name column contents can be alphabetically sorted by clicking on the title header of each cell. This feature is helpful when long list of fluorophores are created as with multiple files (see Gene Study).

=	Setting	Data Table	
€Ge	ene List	Condition List Data Set List	
	Name 🛆	Full Name	Created Date
1	1-Cy5	Sample Gene Expression Time Course3.opd	1/27/2006 5:04:05 PM
2	1-FAM	Sample Gene Expression Time Course3.opd	1/27/2006 5:04:05 PM
3	1-HEX	Sample Gene Expression Time Course3.opd	1/27/2006 5:04:05 PM
4	1-TexasRed	Sample Gene Expression Time Course3.opd	1/27/2006 5:04:05 PM
5	2-Cy5	Sample Gene Expression Time Course2.opd	1/27/2006 5:04:05 PM
6	2-FAM	Sample Gene Expression Time Course2.opd	1/27/2006 5:04:05 PM
7	2-HEX	Sample Gene Expression Time Course2.opd	1/27/2006 5:04:05 PM
8	2-TexasRed	Sample Gene Expression Time Course2.opd	1/27/2006 5:04:05 PM
9	3-Cy5	Sample Gene Expression Time Course1.opd	1/27/2006 5:04:05 PM
10	3-FAM	Sample Gene Expression Time Course1.opd	1/27/2006 5:04:05 PM
11	3-HEX	Sample Gene Expression Time Course1.opd	1/27/2006 5:04:05 PM
12	3-TexasRed	Sample Gene Expression Time Course1.opd	1/27/2006 5:04:05 PM

Fig. 5.43. Assigning and View the Data Set List.

5.9.5 Applying an Analysis Method

Choose an analysis method by clicking one of the option buttons as shown in the screen example below:

- 1. Normalized Expression $(\Delta\Delta C_{T})$ is the default. This option requires you to choose a reference gene.
- 2. Relative Quantity (ΔC_T)

	lormalized e elative quan	xpression (dd tity (dCt)	Ct)		
9	10	11	12		
nown	Unknown	Unknown	Unknov	wn	
beta	Tubulin	Tubulin	Tubulin	1	
23.46	23.25	21.46	19	9.68	
5	*0 Hr	1 Hr	2 Hrs		
nown	Unknown	Unknown	Unknov	wn	

Fig. 5.44. Applying an Analysis Method.

Recalculate to Apply new Settings

Many changes made to the plate setup and analysis options require that you recalculate. If you do not immediately see the change you expect to see, click Recalculate as shown below.

Gene	e Name: IL 1b		▼ Co	ndition Name	Std-7		•	
	Copy cond	ition to all dat	ta sets 💻	Enablefor	Gene Study	E	Analyze V	Vells
	√ ₁	2	3	4	5	6	7	8
			IL1b	IL1b	IL1b	IL1b	IL1b	IL1b
			12.60	15.91	19.23	22.69	26.03	29.42
G			Std-1	Std-2	Std-3	Std-4	Std-5	Std-6
			Standard	Standard	Standard	Standard	Standard	Standard

Fig. 5.45. Updating Gene Expression Analysis Options.

Considerations for Multiplex Analysis

The Copy condition to all data sets button, shown circled in red below, should be active if you want all dye layers or data sets to have the same conditions. This is a useful function if:

- 1. You have a multiplex experiment. In this case all samples in all dye layers must be the same; and
- 2. When multiple OPD (data files) import is enabled; many plate setups will be the exactly the same.

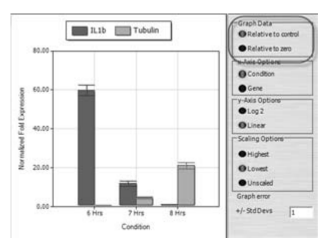


Fig. 5.46. Copying Gene and Condition Settings to all dye layers.

Make sure the button is highlighted (active) when you assign condition (sample) names. The Copy condition to all data sets button is active by default.

Copy Conditions to All Data Sets

The Copy conditions to all data sets button should be activated if you want all dye layers or data sets to have the same conditions. This is a useful function if:

- 1. You have a multiplex experiment. In this case all samples in all dye layers must be the same.
- 2. When multiple OPD (data files) import is enabled in the software many plate setups will be the exactly the same.

Make sure the Copy conditions to all data sets button is highlighted (active) when you assign condition (sample) names as shown below. Active is the default state for this button.

5.9.6 Graphing Options for Expression Data

Graph Expression Relative to Control or Relative to Zero

These options, shown below, allow you to present data with bars originating at 1 (relative to control) or at zero (relative to zero). If you assign a control in your dataset, selecting the option to graph data relative to control allows you to quickly visualize up-regulation and down-regulation. The values graphed are exactly the same.

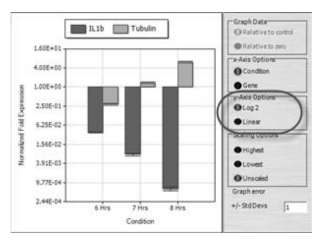


Fig. 5.47. Graphing options for x-axis standards.

Graph y-axis linear or Log2 (Natural Log, Ln)

This option allows you to display the graph with the y-axis in log2 or linear scale as shown below. The log2 scale is useful when you evaluate samples across a large dynamic range.

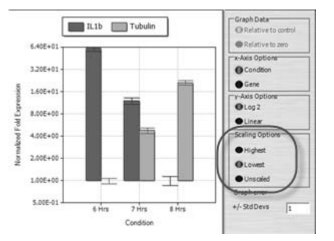


Fig. 5.48. Display options for Gene Expression Graph.

Scaling Options

These options, shown below, are only active in Normalized Expression mode; they allow you to calculate and present your data in a manner that is best suited for your experiment.

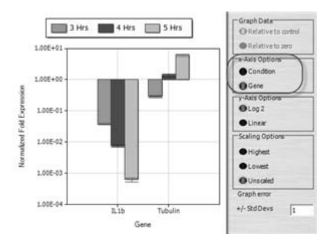


Fig. 5.49. Scaling options for Normalized Gene Expression.

Scale to Highest

This option recalculates the Normalized expression for each gene by dividing the expression level of each condition by the highest expresser. The highest expresser for each gene has a value of 1.

Scale to Lowest

This option recalculates the Normalized expression for each gene by dividing the expression level of each condition by the lowest expresser. The lowest expresser for each gene has a value of 1.

Unscaled

This option does not scale to any sample in particular. It presents unscaled normalized expression.

Scale to Control

Scale to control is another scaling option which is accomplished by assigning a control in the Condition List.

X-axis Options

You may graph either genes or conditions on the x-axis by changing these options.

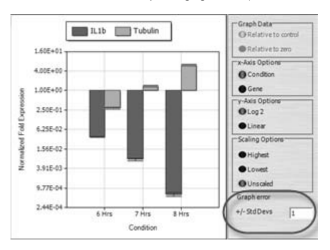


Fig. 5.50. Selecting x-axis grouping options.

Graph Error Options

The default presentation for the error bars is plus and minus one standard deviation. You can change the multiplier to get plus and minus 2 or 3 standard deviations.

	Setting		ata Table								
	DataSet 🛆	Condition 🛆	Gene 🛆	Ctrl	Rel Quant	Rel Quant SD	Corrected Rel Quant SD	Unscaled Expression	Unscaled Expression SD	Corrected Unscaled Expression SD	Expression
1	1-HEX	0 Hrs	Actin	*	1.00000	0.00927	0.00927	N/A	N/A	N/A	N/A
2	1-Cy5	0 Hrs	Gapdh	*	1.00000	0.01676	0.01676	N/A	N/A	N/A	N/A
3	1-TexasRed	0 Hrs	IL1b	*	1.00000	0.02620	0.02620	1.00000	0.02790	0.02790	1.00000
4	1-FAM	0 Hrs	Tubulin	*	1.00000	0.01385	0.01385	1.00000	0.01683	0.01683	1.00000
5	1-HEX	1 Hr	Actin	1	0.99229	0.01295	0.01295	N/A	N/A	N/A	N/A
6	1-Cy5	1 Hr	Gapdh		1.03749	0.03830	0.03830	N/A	N/A	N/A	N/A
7	1-TexasRed	1 Hr	IL1b	1	0.23317	0.01008	0.01016	0.22981	0.01090	0.01098	0.22981
8	1-FAM	1 Hr	Tubulin		3.94348	0.14635	0.15284	3.88658	0.16308	0.16876	3.88658
9	1-HEX	2 Hrs	Actin	1	1.03136	0.01796	0.01796	N/A	N/A	N/A	N/A
10	1-Cy5	2 Hrs	Gapdh		1.11325	0.01443	0.01444	N/A	N/A	N/A	N/A
11	1-TexasRed	2 Hrs	IL1b	1	0.05608	0.00146	0.00158	0.05234	0.00148	0.00158	0.05234
12	1-FAM	2 Hrs	Tubulin		20.49473	2,15090	2.20914	19.12670	2.01803	2.07211	19.12670

Fig. 5.51. Selecting Error Bar options for Gene Expression.

5.9.7 Normalized Expression Calculations

The normalized expression calculation is stated below. To see how the iQ5 software calculates relative quantities, go to Relative Quantity Calculations.

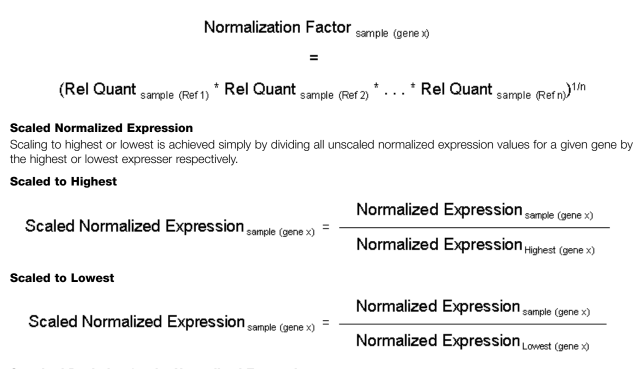
Normalized Expression_{sample (gene x)} = (Rel Quant _{sample (Ref 1)} * Rel Quant _{sample (Ref 2)} * . . . * Rel Quant _{sample (Ref n)})^{1/n}

Normalized Expression when a Control is Chosen

When a control is chosen, the iQ5 software uses the equation listed above to calculate normalized expression. If a control is assigned, relative quantities for all genes within the control sample are equal to 1. This results in "normalized" expression of 1 for the control, relative quantities for all other conditions will be presented relative to this normalized control sample. According to the calculations performed by the iQ5 software, Normalized Expression when a control is chosen is equivalent to unscaled normalized expression analysis.

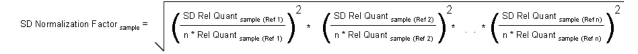
Normalization Factor

The denominator of the normalized expression equation is referred to as the normalization factor. The normalization factor is the geometric mean of the relative quantities of all the reference genes for a given sample.

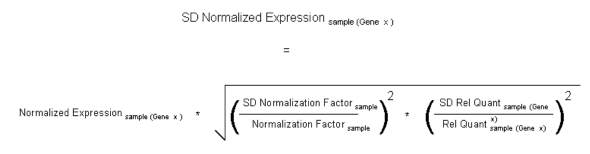


Standard Deviation for the Normalized Expression

Rescaling this value is accomplished by dividing the Standard deviation of the normalized expression by the normalized expression value for the highest or lowest expresser depending on which scaling option you choose.



When a control is assigned you need not perform this rescaling function on the standard deviation as illustrated below.



Standard Deviation of the Scaled Normalized Expression

Rescaling this value is accomplished by dividing the standard deviation of the normalized expression by the normalized expression value for the highest or lowest expresser depending on which scaling option you choose.

SD Scaled Normalized Expression $_{sample (Gene \times)} =$

SD Normalized Expression _{sample (Gene ×)}

Normalized Expression MAX or MIN (Gene x)

When a control is assigned you need not perform this rescaling function on the standard deviation.

5.9.8 Relative Quantity Calculations

Relative Quantity Calculations with No Controls Identified

Relative Quantity (ΔC_T) for any sample for gene x is calculated as follows.

Relative Quantity_{sample (Gene x)} =
$$E_{\text{Gene x}}^{(C_{T(MIN)} - C_{T(sample)})}$$

Where E = Efficiency of primer/(probe) set

this efficiency is calculated as follows (% Efficiency * 0.01 + 1) where 100% = 2

 C_{T} (MIN) = Average C_{T} for the sample with the lowest average C_{T} for Gene x

 C_{T} (sample) = Average C_{T} for the sample.

Relative Quantity When a Control is Assigned

With a control assigned Relative Quantity (ΔC_T) for any sample for all genes is calculated as follows.

$$(C_{T(Control)} - C_{T(sample)})$$
Relative Quantity_{sample (Gene x)} = $E_{Gene x}$

Where E = Efficiency of primer/(probe) set

this efficiency is calculated as follows (% Efficiency * 0.01 + 1) where 100% = 2

 C_{T} (Control) = Average C_{T} for the sample which has been assigned as a control

 C_{T} (Sample) = Average C_{T} for the sample

This is where the calculations differ from those outlined by Dr. Jo Vandesompele on the geNorm Web site. In the example on the geNorm Web site, the results are not scaled the control until normalized expression is calculated. This is referred to as rescaled normalized expression in the example spreadsheet.

Standard Deviation of Relative Quantity of Gene x for a Given Sample

SD Relative Quantity = SD C_T sample * Relative Quantity sample * Ln(E(Gene x))

Where:

SD Relative Quantity = Standard Deviation of the Relative Quantity

SD CT sample = Standard Deviation of the C_{T} of the sample.

Relative Quantity sample = Relative Quantity of sample.

E = Efficiency of primer/(probe) set

this efficiency is calculated as follows (% Efficiency * 0.01 + 1) where 100% = 2

5.9.9 Corrected Values Calculations

The Efficiency value (E) used in gene expression calculations has an associated error. If a standard curve was generated as part of the real time PCR assay, this error can be calculated and used to adjust the error associated with the following standard deviation values:

- Rel Quant SD
- Unscaled Expression SD
- Expression SD

Corrected Values for all gene expression results are displayed on the Data Table spreadsheet when the Show Details button is selected. When Show Details is active, three new columns appear in the Data Table spreadsheet. The new columns display the error correction propagated from the standard curve data for

- Relative Quantitation Standard Deviation
- Unscaled Expression Standard Deviation
- Expression Standard Deviation

The new columns are named:

- Corrected Rel Quant SD
- Corrected Unscaled Expression SD
- Corrected Expression SD

						Rel Quant	Corrected	Unscaled	Unscaled	Corrected Unscaled	
	DataSet 🛆	Condition 🛆	Gene 🛆	Ctrl	Rel Quant	SD	Rel Quant SD	Expression	Expression SD	Expression	Expression
1	1-HEX	0 Hrs	Actin	*	1.00000	0.00927	0.00927	N/A	N/A	N/A	N/A
2	1-Cy5	0 Hrs	Gapdh	*	1.00000	0.01676	0.01676	N/A	N/A	N/A	N/A
3	1-TexasRed	0 Hrs	IL1b	*	1.00000	0.02620	0.02620	1.00000	0.02790	0.02790	1.00000
4	1-FAM	0 Hrs	Tubulin	*	1.00000	0.01385	0.01385	1.00000	0.01683	0.01683	1.00000
5	1-HEX	1 Hr	Actin		0.99229	0.01295	0.01295	N/A	N/A	N/A	N/A
6	1-Cy5	1 Hr	Gapdh		1.03749	0.03830	0.03830	N/A	N/A	N/A	N/A
7	1-TexasRed	1 Hr	IL1b	1	0.23317	0.01008	0.01016	0.22981	0.01090	0.01098	0.22981
8	1-FAM	1 Hr	Tubulin		3.94348	0.14635	0.15284	3.88658	0.16308	0.16876	3.88658
9	1-HEX	2 Hrs	Actin	12	1.03136	0.01796	0.01796	N/A	N/A	N/A	N/A
10	1-Cy5	2 Hrs	Gapdh		1.11325	0.01443	0.01444	N/A	N/A	N/A	N/A
11	1-TexasRed	2 Hrs	IL1b		0.05608	0.00146	0.00158	0.05234	0.00148	0.00158	0.05234
12	1-FAM	2 Hrs	Tubulin		20,49473	2.15090	2,20914	19.12670	2.01803	2.07211	19.12670

5.51 The Gene Expression analysis Data Table - Show Details View.

A difference between corrected values and non-corrected values will only be seen if a standard curve was created as part of the real time PCR assay. The equations used in determining the error propagation are the Standard Error, Standard Error for Normalized Expression and the Standard Error for the Normalized Gene of Interest (GOI).

The equation for Standard Error is shown below:

$$SE = \frac{SD}{\sqrt{n}}$$

Where n = number of measurements

The Standard Error for Normalized Expression equation is shown below:

$$SE NF_{n} = NF_{n} \times \sqrt{\left(\frac{SE \operatorname{Re} l \operatorname{Quant}_{sample (\operatorname{Re} f 1)}}{n \times \operatorname{Re} l \operatorname{Quant}_{sample (\operatorname{Re} f 1)}}\right)^{2} + \left(\frac{SE \operatorname{Re} l \operatorname{Quant}_{sample (\operatorname{Re} f 2)}}{n \times \operatorname{Re} l \operatorname{Quant}_{sample (\operatorname{Re} f 2)}}\right)^{2} + \dots + \left(\frac{SE \operatorname{Re} l \operatorname{Quant}_{sample (\operatorname{Re} f n)}}{n \times \operatorname{Re} l \operatorname{Quant}_{sample (\operatorname{Re} f n)}}\right)^{2}$$

Where NF = Normalization Factor

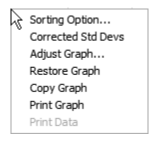
The Standard Error for Normalized GOI equation is the following:

$$SE \quad GOI_{norm} = GOI_{norm} \times \sqrt{\left(\frac{SE \quad NF_n}{NF_n}\right)^2 + \left(\frac{SE \quad GOI}{GOI}\right)^2}$$

Where GOI_{norm} = the Normalized Gene of Interest

5.9.10 Gene Expression Module Context Menus

Right-clicking on the Gene Expression chart will reveal the context menu shown below. The Gene Expression chart context menu contains the following options:



5.52. The Gene Expression chart context menu.

Sorting Option...

The Sorting Option allows for sorting of the Gene Names and Condition Names on the chart display. When the Sorting Option is selected a Sort Options menu box is displayed:

Sort Options		×
Groups Gene Names Condition Names Options Alphabetic order Manual order OK Cancel	Actin Gapdh IL1b Tubulin	

Fig. 5.53. The Sort Options dialog box.

The menu box is divided into three sections: Groups, Options, and the Group member list. Use the **Groups** radio buttons to view a list of Gene Names or Condition Names configured in your data set. With the Options radio buttons, you can elect to organize the names within the Group member list alphabetically, or to rearrange these items manually.

- **Alphabetic order**: The default selection for Groups is Gene Names and the default selection for Options is Alphabetical order. The arrows buttons to the right of the group member list can be used to toggle between an alphabetical sort of the group members in ascending or descending order.
- **Manual order**: To rearrange Gene or Condition Names manually, select the radio button for Manual order. Select the group member that you would like to re-order and use the arrow buttons to move the member to the desired position within the list of names. More than one Gene or Condition Name can be selected (highlighted) by clicking on the desired names. Note that when the Manual order radio button is active, a total of four arrow buttons are displayed. The outermost arrow boxes move the selected items to the top or bottom of the list, while the innermost arrow buttons allow for stepwise movement of the Gene Names or Condition Names.
- When finished sorting, click OK to view the changes to the chart.

Corrected Std Devs

The Corrected Std Devs selection enables the display of error correction propagated from the standard curve data. This option should only be used if a standard curve was created as part of the gene expression analysis. The resultant error correction is displayed as a property of the individual error bars on the chart. The exact error value can be displayed by mousing over the error bars to reveal the tool tip or by displaying the Data Table and then selecting Show Details. When Corrected Std Devs is active, a check mark is displayed next to its name in the Gene Expression chart context menu.

Refer to Section 5.5.2 for details on the Adjust Graph, and Restore Graph options of the chart context menu.

Gene Expression Chart Tool Tips

Tool Tips are visible on the Gene Expression chart when the mouse cursor is positioned on one of the chart data bars or error bars.

When the mouse cursor is positioned on the chart data bars, the resulting Tool Tip will display the data label (i.e., Gene or Condition Name, depending on which x-axis mode is selected) along with the calculated Expression value in parenthesis:

When the mouse cursor is positioned on the chart error bars, the resulting Tool Tip will display the Condition Error or Gene Error, depending on which x-axis mode is selected. The error value displayed is the Unscaled Expression standard deviation. However, the Tool Tip will display the Corrected Unscaled Expression standard deviation if Corrected Std Devs is selected from the Gene Expression context menu.

Copy Graph / Export a Graph

To export a graph:

- 1. Place the mouse pointer over the graph.
- 2. Click the right mouse button. A shortcut menu appears.
- 3. Click Copy Graph.
- 4. Switch to or open the document into which you will paste the graph.
- 5. Paste the graph image by choosing Edit, then Paste, or by pressing CTRL+V.

Export Data from the Data Table

- 1. Make sure the Data Table button is activated. If not, click Data Table.
- 2. Click the right mouse button. A shortcut menu appears. See the screen example below.
- 3. Select Export to Excel.
- 4. Enter a name and file destination for the Excel file generated with your data export.

-	Data Table
\mathbf{k}	Print
ľ	Export to Excel

Fig. 5.54. Exporting Gene Expression data tables to Microsoft Excel.

The Print command, on the Spreadsheet menu, will print the displayed Spreadsheet (Gene List, Condition List, Data Set List or Data Table). When selected, a Print Preview box is opened which contains an illustration of the spreadsheet as it will appear once printed. Clicking the printer icon opens the Windows print dialog box. Click OK to complete the printing task.

The Export to Excel command on the Spreadsheet menu is useful for exporting exact values from the spreadsheet. When the Export to Excel command is selected from the menu, an Export to Excel file save box is displayed. Choose a location of where the Excel file is to be saved and click Save. The iQ5 software automatically exports the selected data into a protected workbook.

- The protected workbook generated by the iQ5 software contains the text values of what is represented on the Spreadsheet. For example, check boxes from the software application are replaced by "True" or "False" text in Excel.
- The numeric values contained in the protected workbook are exact values from the software application that include several non-significant figures beyond the decimal point. This is important to note when considering whether to transfer iQ5 spreadsheet data by a copy and paste command or the Export to Excel command. With the copy and paste command only the significant digits displayed in the iQ5 software interface are transferred to Excel.

Gene Expression Plate Interface Context Menu

The Plate Interface menu is displayed by positioning the cursor over the Plate Interface and clicking the right-mouse button. When this action is performed, a menu is displayed with the following items: Copy Gene Name, Copy Condition Name, Paste Gene/Condition Name, Show Sample Type, Print and Export to Excel.

	1	2	3	4	5	6	7	8	9	10	11	12
А												
в				19.93 *0 Hrs	17.79 1 Hr	Tubulin 15.46 2 Hrs Unknown						
с				19.89 *0 Hrs	17.85 1 Hr	Tubulin 15.20 2 Hrs Unknown			Copy C Paste G	ene Name ondition Nam iene/Conditio ample Type		
D				19.90 *0 Hrs	17.90 1 Hr	Tubulin 15.49 2 Hrs Unknown			Print	to Excel		

Fig. 5.55. The context menu of the Gene Expression Plate Interface.

- Copy Gene Name and Copy Condition Name: These two commands are useful shortcuts for copying and pasting Gene Name and Condition Name information across many different samples within a single .opd or Gene Study.
 - 1. Activate the copy command by first selecting the wells to be copied from the Plate Interface.
 - 2. Right-click on the plate interface to reveal the Copy Gene or Copy Condition options, and select the desired option.
 - 3. Once the Copy Gene or Copy Condition option has been selected, the Paste Gene/Condition Name text from the menu becomes active (no longer grayed-out).
 - 4. Return to the Plate Interface to select the destination wells for the copied identification information.
 - 5. Right-click on the plate interface to reveal and select the Paste Gene/Condition Name option.
- Show Sample Type: The Show Sample Type option is selected by default. This selection is illustrated by a check mark shown before the text in the menu. Clicking on Show Sample Type toggles the display of the Sample Type in the Plate Interface. Choosing to remove the sample type information from the Plate Interface simply allows visibility of a greater number of wells in the Plate Interface grid.
- Print: The Print command, on the Plate Interface menu, will print the displayed Plate Interface.

• **Export to Excel**: The Export to Excel command on the Plate Interface menu is useful for exporting exact values from the Plate Interface. When the Export to Excel command is selected from the menu, the iQ5 software exports the selected data into a protected workbook that is automatically open by Microsoft Excel. The protected workbook contains all of the formatting and fluorophore information separated into worksheets.

Note: The numeric values contained in the protected workbook are exact values from the software application that include several non-significant figures beyond the decimal point. This is important to note when considering whether to transfer iQ5 spreadsheet data by a copy and paste command or the Export to Excel command. With the copy and paste command only the significant digits displayed in the iQ5 software interface are transferred to Excel.

5.9.11 Reaction Efficiency and Gene Expression Analysis

Efficiency describes how much of your sequence of interest is being produced with each cycle. 100% efficiency means that you are doubling your sequence of interest with each cycle. People focus on efficiency for a number of reasons. There is evidence that using accurate efficiencies for each of your primer/(probe) pairs will give you more accurate results when using the mathematical modules used in the Gene Expression tab.

You set the efficiency for each of your genes (sequences, primer/(probe) set) in the Gene List. The default value is 100 percent efficiency.

Auto Efficiency

The Auto Efficiency function uses efficiency values from your experiment so that you do not have to type them in yourself. To use Auto Efficiency, you must have standards in your experiment that result in valid standard curves in the PCR Quant tab.

Auto Efficiency Standards

The iQ5 software only requires two standards at different concentrations though it is recommended to have at least four samples in triplicate across a relevant dynamic range.

5.9.12 Gene Expression – Frequently Asked Questions

Why should I normalize my data?

Relative Quantity data that is not normalized by some means is difficult to interpret. Imagine the case where you load 1 µg of RNA in one well and 10 ng in the other well. If you perform a relative quantity analysis on the results from such an assay the fact that the 10 ng sample has a smaller Relative Quantity value is irrelevant. It is likely the result of using less RNA and not the result of some biological response.

How does normalized expression as calculated by this software compare to the $\Delta\!\Delta C_{_T}$ equation?

If you leave efficiencies at 100 percent and only evaluate one reference gene and one gene of interest the software will generate the same results as you would get using the $\Delta\Delta C_{\tau}$ equation.

The standard deviations will be larger since the error propagation outlined in the initial publication is inappropriate.

How does normalized expression as calculated by this software compare to the model introduced by Dr. M. Pfaffl, et al.?

If you only evaluate one reference gene and one gene of interest you will get exactly the same results using the iQ5 software as you would using the model introduced in this paper.

Standard deviations may be slightly different.

How does normalized expression as calculated by this software compare to the model outlined by Dr. Jo Vandesompele on the geNorm Web site?

The iQ5 software uses the models outline on the geNorm Web site and will give you the same results.

Why would I have to assign gene names in the gene expression tab?

If you have not assigned gene names in your initial plate setup or if you are studying more than 5 genes you can use the plate interface in the gene expression tab to assign these relationships.

Can I customize my gene and condition names?

Yes. Both the gene pull-down menu and the condition pull-down menu will accept any text you type into them. If your names are very long, you can edit the long name field in the gene list and condition list. The long name will only be displayed in the legend above the graph. The name displays on the x axis.

How do I Determine Efficiencies?

Typically the efficiency for each primer (or primer/probe) set is evaluated and recorded during assay development. Generate a standard curve using serial dilutions of a representative sample across a relevant dynamic range, and then record the efficiency for subsequent gene expression analysis

5.10 Gene Study: Multi-File Gene Expression Analysis

When conducting Gene Expression experiments, it may become necessary to run more than a single plate in order to analyze all required samples. Similarly, the experimental goal may be to analyze samples/conditions over a fixed time course. In both cases, it is essential to perform Gene Expression analysis on data generated in different data sets. To accommodate this need, a Gene Study can be created in the iQ5 software.

5.10.1 The Gene Study File

A Gene Study is a specialized file consisting of data and sample information imported from multiple .opd files. The imported data is grouped into a single study which can be edited (files added or removed) and further analyzed at the users discretion. Gene Study files are assigned a file extension of .gxd by the iQ5 software.

A Gene Study is capable of comparing approximately 5,000 total wells of data. This implies a maximum comparison of approximately 52 .opd files containing 96 wells of single-fluor (non-multiplexed) data, or 10 .opd files containing 96 wells of five-fluor (multiplexed) data. Many other combinations of file number, well capacity, and fluor number are possible. The absolute well maximum will depend on the amount of RAM and virtual memory available to your computer.

Enable for Gene Study Button

All .opd files must be enabled for Gene Study analysis from within the Data Analysis module of the iQ5 software. Enabling an .opd file for Gene Study analysis allows the iQ5 software to extract only the information critical to Gene Expression analysis between different files. This step is performed by activating the Enable for Gene Study button found in the Gene Expression window of the Data Analysis module.

+	Gene	Name:		▼ Cor	ndition Name	:
		Copy cond	ition to all dat	ta sets	Enablefor	Gene Study
		1	2	3	4	5
	А					
		[beta-actin]	[beta-actin]	[beta-actin]		

Fig. 5.56. The Edit Plate sample type icons.

When a file has been saved with the Enable for Gene Study button active, it allows the saved file to be added to a Gene Study. The default setting is that this button is not enabled. You must save your data file after activating this option from within the Gene Expression analysis window.

5.10.2 Creating a Gene Study

A Gene Study must contain data from at least two different OPD files. The process of creating a Gene Study is a 2-step process.

STEP 1: Prepare .opd files for Gene Expression analysis.

a. In the PCR Quant tab, establish the desired experimental conditions such as baseline and threshold and which wells are to be included in the analysis.

Note: Additional wells can be excluded or re-included later from the Gene Expression Plate Interface.

- b. Click on the Gene Expr tab. Make sure that all files to be included in the Gene Study have the **Enable for Gene Study** button selected. The default setting is that this button is not enabled.
- c. After enabling the file for multi-file analysis, save the OPD file.
- d. Repeat steps a-c, above, for all additional files to be included in the Gene Study.

STEP 2: Add files to a new Gene Study.

- a. From the menu toolbar select File/New/Gene Study. The Gene Expression Study Manager will be displayed.
- b. Select Add OPDs. The Windows file explorer dialog box will appear.
- c. Locate and select the OPDs needed for the Gene Study. Multiple files may be added to the Gene Study at one time.
- d. Click OK when finished.

When the files are added to the Gene Expression Study Manager, the file information is separated into four columns:

- File Selection: This column is used in conjunction with the Remove button, and features a check box in the column header. The "header" check box will select or deselect all the displayed files chosen for the Gene Study. To select or deselect an individual file, click the check box within the corresponding row. If one or more files have been selected, clicking the Remove button will remove those file(s) from the Gene Study.
- File Name: This column displays the full name of the .opd files to be included in the Gene Study.
- File Directory: This column displays the directory location of the .opd files to be included in the Gene Study.
- **Created Date**: This column displays the creation dates of the .opd files to be included in the Gene Study. Note that these dates are NOT the dates of the most recent save event for the listed files.

Gene Study Errors

The file must be saved with the Enable for Gene Study button selected. If a file with the Enable for Gene Study is not selected and there is an attempt to add the file to a Gene Study an error will be received:

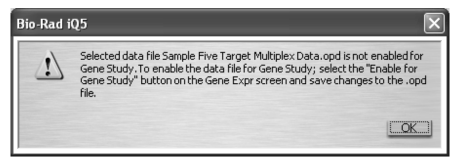


Fig. 5.57. Gene Study error message - file not enabled.

To correct this error:

- 1. Click OK to clear the error message.
- 2. Re-open the file that will be added to the Gene Study.
- 3. Confirm that all wells to be analyzed have valid C_{τ} values and activate the Enable for Gene Study button.
- 4. Save the file.
- 5. Attempt to add that file to the Gene Study once again.

5.10.3 Gene Expression Study Manager

The columns and rows of the Gene Expression Study Manager are adjustable by clicking, holding and dragging the column/row separator lines. The overall size of the Gene Expression Study Manager box can also be resized by clicking, holding and dragging the outermost horizontal or vertical edge.

The Notes section, located at the bottom of the Gene Expression Study Manager, allows for input of text details pertaining to the study.

Clicking Show Details, located below the OPD spreadsheet, displays two additional columns of information. The two additional columns provide information about the Cycle and Step used for data collection and analysis in the source .opd file.

Add		xpression Study Manager move OPD files						
		File Name		File Directory	Created Date	Cycle	Step	
1		Sample Gene Expression Time Course3	H:\		3/14/2006 4:56:11 PM	2	2	1
2		Sample Gene Expression Time Course2	H:\		3/14/2006 4:56:11 PM	2	2	
3		Sample Gene Expression Time Course1	H:\		3/14/2006 4:56:11 PM	2	2	1
							1.	
•	۸dd O	PDs Remove			Show Details			
-		PDs Remove Multiplex Gene Study Example			Show Details			28-80.00
4			хп		Show Details			

Fig. 5.58. The Gene Expression Study Manager -Show Details View

Once the files are added to the Gene Expression Study Manger, selecting OK opens an unsaved Gene Study file in the Gene Expr window of the iQ5 software. The Gene Study, which contains all the files selected from the Gene Expression Study Manager, can now be analyzed as a normal Gene Expression file by selecting the analysis method, assigning Gene and Condition Names from the Gene Expression Plate Interface and assigning attributes from the Gene List.

Note: Gene Study files are **NOT** automatically saved at the time of file creation. It is highly recommended to save the newly created .gxd file after initiating a Gene Study.

5.10.4 Editing a Gene Study

A minimum of two different .opd files are required to create a new Gene Study file. Additional .opd files may be added to an existing Gene Study file by clicking the Edit Study button from within the Gene Expr window. The Edit Study button is only displayed in the Gene Expr window when a Gene Study file is being analyzed.

When you click on the Edit Study button, the iQ5 software will display the Gene Expression Study Manager. Individual files can be added to the Gene Study by clicking Add OPDs. To remove files from the current Gene Study, To select or deselect an individual file, click the check box within the corresponding row. If one or more files have been selected, clicking the Remove button will remove those file(s) from the Gene Study.

The Notes field is also editable from the Gene Expression Study Manager. Once the desired changes have been made, click OK to close the Gene Expression Study Manager and return to analysis of the Gene Study file in the Gene Expr module. Remember to immediately save your edits to the Gene Study before continuing with your analysis.

Note: The Edit Plate module of the iQ5 software is not available when a Gene Study file (extension, .gxd) is opened. All plate information changes to the data in a Gene Study(i.e. sample ID), must be made from the Plate Interface of the Gene Expr window.

5.10.5 Managing Samples from the Gene Expression Plate Interface

When a Gene Study file is displayed, the Analyze Wells button that is available when viewing a single Gene Expression file is no longer displayed. In its place is the Include Sample/Exclude Sample drop down list located above the Plate Interface.

Include Sample	•
Include Sample	
Include Sample Exclude Sample	

Fig. 5.59. The Include Sample / Exclude Sample selection box.

This drop down list works similar to Analyze Wells button (see Section 5.2.4 for details). To exclude wells from the Gene Study analysis, select the specific wells from the Plate Interface. Click on the drop down arrow box to select Exclude Sample from the list. The highlighted wells will be grayed to indicate that they are no longer included in the Gene Study analysis. To include wells for the Gene Study analysis, select the specific wells from the Plate Interface. Click on the drop down arrow box to select Include Sample from the list. The selected wells for the Gene Study analysis, select the specific wells will be displayed with the correct Gene and Condition Name colors to indicate that they are included in the Gene Study analysis.

5.10.6 Inter-run Calibration

In a Gene Study, data for a given gene may be spread over multiple runs (.opd files). The data from these multiple runs should only be compared in a Gene Study after the data from the runs have been calibrated to each other using the Inter-run calibration algorithm.

An Inter-run calibrator is an identical sample (i.e. primer/template or primer/probe/template combination) that is present on every plate in the Gene Study. For a given gene, it is assumed that differences in C_{τ} values between plates for identical samples are due to inter-run variations. Therefore, the C_{τ} value differences between identical samples can be used as "Inter-run calibrators" for a given gene. Inter-run calibrators are used to counteract the effects of inter-run variation from all other samples of a given gene. Ideally, more than one inter-run calibrator would be run on every plate.

Note: Genes which have no samples that can be used for Inter-run calibration will be processed without correction in the Gene Study (not recommended).

Inter-run Calibration is unique to the Gene Study module. Inter-run Calibration is automatically attempted by the iQ5 software in every Gene Study. The iQ5 software identifies an inter-run calibrator as a sample with the exact same Condition Name on each plate (per assay) and for each gene – this allows the sample to be designated as an "inter-run calibrator". The resulting Inter-run Calibration calculations are then used to normalize inter-run variations between genes assayed in separate real-time PCR runs (i.e., different .opd files). The following conditions must be met for Inter-run Calibration to be performed:

- A given sample must have the exact same Condition Name on each plate (per assay) and for each gene this allows the sample to be designated as an "inter-run calibrator".
- At least one "inter-run calibrator" sample must be present in the Gene Study.

When the above conditions are met, an algorithm is used to calculate the pair-wise differences between the C_{τ} values (ΔC_{τ}) for all samples that qualify as inter-run calibrators (i.e., having the same Condition Name per gene and per assay).

Inter-run Calibration Algorithm

All data within the Gene Study is normalized with to the inter-run calibrator that yields the smallest average ΔC_{τ} value. The inter-run calibrator with the smallest average ΔC_{τ} value becomes the dominant inter-run calibrator. The average ΔC_{τ} value of the dominant inter-run calibrator will be used to adjust all C_{τ} values within the Gene Study.

To find the dominant inter-run calibrator, the iQ5 Inter-run Calibration algorithm first calculates the average of the ΔC_{τ} values for all inter-run calibrators of a given gene. The iQ5 software uses a multi-tiered algorithm to determine the dominant inter-run calibrator.

The algorithm utilizes the following hierarchy for determining the dominant inter-run calibrator:

1. Set the dominant calibrator to the gene with the highest number of common replicate groups in a given pair-wise comparison.

If multiple genes have the same number of common replicate groups, then:

2. Set the dominant calibrator to the gene with the smallest range of ΔC_{τ} values in pair-wise comparisons. The range is examined by comparing the absolute value of the difference between the maximum and minimum ΔC_{τ} for the inter-run calibrators of a given gene.

If multiple genes have an identical range of ΔC_{τ} values, then:

3. Set the dominant calibrator to the gene with the smallest absolute value of "Average ΔC_{τ} " of eligible inter-run calibrator samples.

If multiple genes have identical "Average ΔC_{τ} " absolute values, then:

4. Set the dominant calibrator to the replicate group with the smallest ΔC_{τ} .

Note: The first data file imported into the Gene Study will always serve as the "hub" file for pair-wise data comparisons during Inter-run Calibration.

Se	lect gene :	Actin	•		
	Inter-run calibrator	1-HEX	2-HEX	dCt	
1	Std-3	18.2092	18.2156	-0.0064	
2	Std-5	25.7935	25.3787	0.4148	
3	Std-4	22.5125	22.5923	-0.0798	
4	Std-6	29.5600	29.7665	-0.2065	_
5	Std-7	33.4224	33.3596	0.0628	
6	Std-1	12.6251	12.5211	0.1040	
7	Std-2	15.1694	15.5875	-0.4181	
8			Average dCt	-0.0185	
9			ddCt	0.8329	
M	▲ ▶ ▶ 1-HEX \	/s 2-HEX / 1-HEX vs	3-HEX /	•	Þ

Fig. 5.60. The Inter-run Calibrator calculation display.

Details of the Inter-run Calibration calculations can be accessed by clicking the Inter-run Calibration button. The resulting Inter-run Calibration window (shown above) will display the comparative fluorophore calculations per gene (pair-wise comparisons).

- You can choose to view the inter-run calibrator calculations for different genes in your assay by selecting a Gene Name from the Select Gene drop-down menu.
- You can choose to view the calculations resulting from different pair-wise comparisons by clicking on the desired tab displayed below the spreadsheet.

Performing a Gene Study without Inter-run Calibration

When data files are merged into a Gene Study that does not meet the criteria for Inter-run calibration, a warning message will be displayed. The warning message states that the genes being analyzed do not contain common samples across all data sets. The warning message will also list all of the genes in the Gene Study that do not have samples eligible for Inter-run calibration. By clicking the OK button of the warning message, the analysis will proceed without any inter-run calibrators.

Clicking on the Inter-run Calibration button when there are no calibrators assigned will display a message box which states that inter-run calibrators are not available. Proceeding without inter-run calibrators may cause inaccurate representation of the data. To correct this problem change the Gene and/or Condition Names in the Plate Interface so that identical Gene and Condition Names are used across all data sets of the Gene Study. Use the Gene Name and Condition Name drop down menus to select the appropriate Gene and Condition Names or directly input the correct Gene and Condition Names. To change a large number of Gene and Condition well names across several plates use the Copy and Paste command from the right mouse button menu.

5.11 Edit Plate

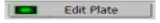
The Post-Run Edit Plate window allows you to edit the plate setup of a data file after an experiment has been performed. This feature allows you to make corrections for incorrect sample type, identifiers/conditions, probe/primer names, units, or standard concentration assignments and to add notes about the experiment.

You may not add or remove fluorophores from wells in post-run plate editing, nor may you delete a well from the plate setup by removing both its sample type and fluorophore assignments. (Use Analyze Wells to remove undesired wells from the analysis.) After editing the plate, you can reanalyze the experimental data with the new plate setup.

Note: You can always restore the original plate setup definitions by clicking Restore Original Plate.

For post-run editing of the Plate Setup saved in a data file:

- 1. From the Workshop module:
 - a. Click Data File above the directory of the home workshop. Navigate the directory until the desired data file is found. Double-click the file name to bring the file directly into the Data Analysis module.
 - or
 - b. Click Data File above the directory of the home workshop. Navigate the directory until the desired data file is found. Click the file name once to open the plate setup associated with the data file in the bottom right pane of the Workshop window. Click Analyze to bring the data to the Data Analysis module.
- 2. At the top of the Data Analysis window, click Edit Plate.



3. A modified version of the Plate Setup Editor will open. In this modified window, you cannot add fluorophores to or remove fluorophores from the fluorophore list. Nor may you add a previously defined fluorophore or remove a previously defined fluorophore from a well.

PCR PCR	Quan	t		Melt Cu	irve/Pe	eak		En	d Poin	t		All	elic Dis	c 🗖 🗖	Gene Expr		Edit Pla
																Restor	e Original Pl
Editing Pla	te :	s	ample G	ene Expi	ession.	pts					-	5ample	Volume	25	÷	Apply	Plate Chanc
Notes :												5eal Ty	pe:	Film	-	Can	cel Change
		ļ									<u> </u>	/essel 1	fype :	Plates	~		
Spreadshe	eet	N	S	x		NTC	()			R				Experiment Na	me:		nent Type :
Replicates														My experiment.		General	
-		1	2	3	4	5	6	7	8	9	10	11	12	Fluorophore	Probe/Pr	imer	
	А													O FAM	Tubulin		
123		<u> </u>							<u> </u>					•HEX	Actin		
-	В					2	3							OTexasRed	IL1b		
	С					2	3							Cy5	Gapdh	_	
1 2 3		<u> </u>			嵩	H											
Size:	D				1	2	3										
1 ÷	E													Whole Pl	late loading		
Next #:	F				2	3	4	5	6	$\widehat{}$				Units:	copy num	iber	
	-			\ge	X		X	X	X	X				Apply units to:	Whole	Plate	Fluoroph
	G				2	3	4	5	6	\mathcal{O}							
	н				2	3	\odot	5	6	\odot				Scientifi	c Notation		ilution Seri
Rov	v	Colum	ו [Sample	Туре		Rep #				er/Cond	ition			antity		Units
В		4	_	Unknow		~	1) Hrs				N/A		copy nun
			_	Unknow		×	1				D Hrs				N/A		copy nun
			_	Unknow Unknow		×	1				D Hrs D Hrs				N/A N/A		copy num
				OHKNOW	<i></i>	×					JHIS				NVA		copy hu

Fig. 5.62. Editing sample properties with the Edit Plate option.

- 4. Edit any notes about the plate setup in the Notes box.
- 5. Edit the name of the experiment in the Experiment Name box.

- 6. Deselect the Whole Plate loading option button if it is not applicable. Otherwise, leave this option button selected. If Whole Plate loading is unavailable when you begin editing the plate, you may not restore Whole Plate loading.
- 7. To edit a sample type:
 - a. Click a Fluorophore.
 - b. Click the correct sample type icon.
 - c. Select the Next # check box to be sure that the desired number will be assigned to the next standard you define.



- d. Select the type of Replicate loading desired.
- e. Click a previously defined well to overwrite its original sample type definition. Replicates of the well are not affected.
- f. Repeat steps b-e above for each additional sample type to be edited for this fluorophore.
- g. Repeat steps a-f above for each additional fluorophore.
- 8. To edit the concentrations of the standards:
 - a. Click the fluorophore for the standards to be defined.
 - b. Click Dilution Series and calculate the concentration, or enter it directly in the spreadsheet.
- 9. Select the Units for the standards.
- 10. Click Apply Plate Changes to make the changes. To see the effect on analysis, go to one of the other Data Analysis windows.

Apply Plate Changes

11. The original plate setup is retained with the data file and may be restored at any time by clicking Restore Original Plate.

Restore Original Plate

5.11.1 Edit Plate Sample Type Icons

There are seven icon buttons running across the top of the representation of the experimental plate as shown below. These buttons are used to provide the two pieces of information required for each well: sample type and fluorophore(s) to be monitored. The active button is always surrounded by a red box.



Fig. 5.63. The Edit Plate sample type icons.

Beginning at the left, the arrow is the standard pointer. The next five icons define the type of sample in the well: Standard, Unknown, No-Template Controls, Positive Control, and Negative Control, respectively. The paint bucket icon, used to load fluorophores into wells, is not active.

5.12 The Reports Tool

The Reports tool of the iQ5 software is accessible from the menu bar of the iQ5 software. Clicking on the heading for Reports will open the Report Viewer, shown below.

The Reports Viewer pulls data, charts, and graphs directly from the displayed settings of the Data Analysis module currently in use. For this reason, it is best to complete all desired analysis and formatting of the displayed data prior to using the Reports tool.

Report Viewer		
PCR Quantification Repor Select Report:	t	BIO RAD
PCR Quant Detailed	•	1
Sort Data By:		PCR
Replicate #	•	<u>General Data</u>
 Ascending Order Descending Order 		Data File Name Data File Path Collected Data Current Date Run Date User aborted the run
Page Setup		Active RMEs Active Well Factors Background Readings RMF Valid
Print Preview		Well Factors Valid Plate Setup File Name Plate Setup File Path
Print		Protocol File Path Protocol File Path
Save to File		Computer name Created by app Created by user Creation Date

Fig. 5.64. The Report Viewer window for creating new reports.

5.12.1 Report Viewer Options

The following options are available from within the Report Viewer window:

- **Select Report**: Use this drop-down menu to select the level of detail in the report for your dataset. The number and type of reports available will vary according to the Data Analysis Module currently being used. For additional details about the report types, refer to section 5.12.2.
- **Sort Data By**: Within a given report, all data displayed in tables can be sorted by the parameters listed in the drop-down menu. The Sort Data By drop-down menu works in conjunction with **Ascending Order** and **Descending Order** radio buttons. Each of the parameters in the report can be sorted in ascending or descending order by the appropriate radio button. The results of the sort are immediately displayed in the report display area.
- **Page Setup**: Clicking the Page Setup button opens the Windows Page Setup dialog where different page display parameters can be adjusted.
- **Print Preview**: Clicking the Print Preview button opens the Windows Print Preview screen where the printed layout of the pending report can be previewed.
- **Print**: Clicking the Print button opens the Print screen where different print parameters can be adjusted and the report printed.

Save to File: Clicking the Save to File button opens the Save report screen where a specific location can be chosen to save the report in rich text format.

5.12.2 Data Analysis Report Types

Every report template available in the iQ5 software contains a preset report header. This report header outlines very specific information about the dataset being analyzed, such as time and date of file creation, and file path information. At the bottom of the "General Data" section, the iQ5 software provides important information that can be used to determine whether the report contains saved or unsaved changes to data analysis settings.

Report differs from last save

The "Report differs from last save" report entry can be used to determine whether the analysis conditions displayed in the report template are different from the analysis conditions present at the time of the most recent file save event for a given dataset. If this entry reads "Yes", then the displayed data was generated from an unsaved version of the dataset being analyzed.

Notes

Any text entered in the Notes field at the time of a data file creation, or data file save event is included in the header section of every report template.

Protocol

The header of each report template contains a summary of the protocol used for thermal cycling and data collection within a given dataset. Any modifications to the protocol used for data collection will be displayed in the Report Header.

There are over 15 different report templates available in the iQ5 software. The number and type of reports available at any given time will vary according to the Data Analysis module currently being used. The various report options available for each of the Data Analysis modules are summarized below.

PCR Quant Reports

- **PCR Quant Data**: This report template includes only the data included in the Standard Curve/Ct Results spreadsheet the PCR Amplification chart is NOT displayed.
- **PCR Quant Detailed**: This report template includes all of the data available in the PCR Quant module of the iQ5 software including the PCR Amplification chart, Ct value spreadsheet, and analysis parameters. If the dataset being analyzed also contains a standard curve, the standard curve chart and spreadsheet data will also be included in this report.
- **PCR Quant Graph Only**: This report template displays only the PCR Amplification chart, exactly as analyzed and formatted within the PCR Quant module.
- Std Curve Data Only Landscape: This report template includes only the data included in the Standard Curve/Ct Results spreadsheet. Although the report on the screen in displayed in the Portrait orientation, the Page Layout print settings are pre-set to the Landscape Orientation.

Melt Curve Reports

- **Melt Curve Detailed**: This report template includes all of the data available in the Melt Curve/Peak module of the iQ5 software including the Melt Curve chart, Melt Peak Chart, Melt Curve data spreadsheet, and analysis parameters. In addition, if all fluorophores have been selected for display in the PCR Quant module, the PCR Amplification chart and analysis parameters will also be displayed.
- **Melt Curve Graph Only**: This report template displays only the graphs available in the Melt Curve/Peak module of the iQ5 software, such as the Melt Curve chart and Melt Peak Chart, exactly as analyzed and formatted within the Melt Curve/Peak module.
- **Melt Curve Only**: When creating a report from a data set performed as an Melt Curve Only run, the Melt Curve Only report option becomes available. This report template includes all of the data available in the Melt Curve/Peak module of the iQ5 software including the Melt Curve chart, Melt Peak Chart, Melt Curve data spreadsheet, and analysis parameters.

End Point Reports

• End Point detailed: When creating a report from a data set that includes PCR Quantification data, End Point detailed is the only report option available. This report includes a section titled "Data Analysis Parameters", which details the PCR analysis settings for the current dataset.

• End Point only: When creating a report from a data set performed as an End Point Only run, the End Point Only report is the only report option available. This report template contains only the End Point Analysis spreadsheet and data analysis parameters.

Allelic Disc Reports

- Allelic Data Only: The Allelic Data Only report template is limited to the data displayed in the Allelic Disc module only, but will not display the Allelic Discrimination chart.
- Allelic Detailed: This report template includes all of the data available in the Allelic Disc module of the iQ5 software including the Allelic Discrimination chart, spreadsheet, and analysis parameters. In addition, if all fluorophores have been selected for display in the PCR Quant module, the PCR Amplification chart and analysis parameters will also be displayed.
- Allelic Only: This report template is limited to the data displayed in the Allelic Disc module only this includes the Allelic Discrimination chart, spreadsheet, and analysis parameters.

Gene Expression Reports

- Gene Expression Detailed: This report template includes all the Gene Expression data available in the Gene Expr module – including charts, data tables, Gene List and Condition List information, Threshold Crossing Spreadsheet, and Gene Expr analysis parameters. In addition, if all fluorophores have been selected for display in the PCR Quant module, the PCR Amplification Chart and PCR Baseline Analysis Parameters will also be displayed.
- **Gene Expression Graph only**: This report template displays only the Gene Expression chart, exactly as analyzed and formatted within the Gene Expr module.
- **Gene Expression only**: This report template includes all of the data tables, Gene List and Condition List information, Threshold Crossing Spreadsheet, and Gene Expr analysis parameters available in the Gene Expr module the Gene Expression chart is NOT displayed.

Gene Study Report

• **MultiFiles Gene Expression Detailed**: When creating a report from within a Gene Study, "MultiFiles Gene Expression Detailed" is the only selection available. This report format is similar to the Gene Expression Detailed report; however, no PCR Quant data is available for this report

Section 6 Calibration Module

Every experiment on the iQ5 or MyiQ system requires that the instrument be calibrated. You calibrate the iQ5 and MyiQ cameras using the iQ5 software Calibration Module.

There are four calibrations required for the iQ5 device which must be done in the following order:

- Mask Alignment
- Background Calibration
- Persistent Well Factor Generation
- Pure Dye Calibration

There are three calibrations required for the MyiQ device, which must be done in the following order:

- Mask Alignment
- Background Calibration
- Persistent Well Factor Generations

Note: Pure Dye Calibration is NOT performed on the MyiQ real time PCR detection system.

This order is the order that the Calibration tabs occur in the iQ5 Software (from left to right). Choose the Mask Alignment tab first. When you complete each calibration choose the tab to the right of the completed calibration to do the next calibration.

Filter Position	Image										Up
• Blank										Left	
•2			a a				0	-		10.2	Down
01								0	•	Ontin	Nze Mask + -
•							0	6	6		
	10-1	0.0					6	6	0		
•5	(0.1	0.0	0.0				0	•		-	Show Mask
•6			9			•	0	•	6	-	WF correction
Home		. 0	0 0				0	0	0)		Save Mask
Exposure Time (ms)						•	40	40			Reload Mask
512 .											Stranger 2001-0
Take an Exposure										Re	store Factory Mask
			_	_	_	_	_	_	_		
rel France	Camera Status :									Serial Number	EP0009
the second se	Carrens connected	TES								FIV version	1.028
adiground		and second a								Library version	1.824
eckground Fick	Optical lid sensor : O									Model	
eckground Fick verage Imer uorescence Fick	Concelle service : C										DQSCOLOR
adiground PUA										Lamp Time	IQSCOLOR

Fig. 6.1. The Mask Alignment Window.

6.1 The Image Screen

6.1.1 Filter Position

Filter Position
• Blank
•2
03
•4
•5
05
Home
Exposure Time (ms)
512 •
Take an Exposure

Fig. 6.2 The Filter Selection Frame.

The exposure time can be changed from this screen and an exposure taken.

6.1.2 Camera Status

Camera Status :	Serial Number	EP0009
Camera connected 11YES	FW version	1.028
Optical lid sensor : CLOSED	Library version	1.024
	Model	IQSCOLOR
	Lamp Time	5583h 21min
	Warmup Time	\$47h-\$3min

Fig. 6.3. The Camera Status Screen.

The Camera Status screen provides feedback about the iQ5 Camera. In this screen you can determine if

- 1. The camera is connected
- 2. The optical lid is closed
- 3. The serial number, firmware version, library version and model of the camera
- 4. The length of time the camera has been in use since it was installed
- 5. The length of time the lamp has been on since the camera was last turned on.

Calibration factors are specific for the optical path you are using which includes the Reaction vessel, sealing mechanism and volume used in the experiment you wish to run.

You need the following to calibrate the iQ5 Camera:

- iCycler iQ Calibrator Dye Solution Set
- iCycler iQ External Well Factor Solution (found in the iQ Calibrator Dye Solution Set)
- 3 x 96-well PCR plate or preferred reaction vessel
- Optical quality sealing tape or preferred sealing method

With these materials, you must prepare three plates: the external well factor plate, the background calibration plate, and the pure dye calibration plate.

6.2 Mask Calibration



Fig. 6.4. Camera Image with Overlying Mask Array.

The Image screen displays the fluorescence detected by the CCD camera for each of the 96 wells on the plate. Pixels that are saturated will be shown in pink. The Mask is displayed as an array of green boxes. Each individual box should be centered over the well so that all the fluorescent signal from the well falls within the box.

The Mask can be displayed or hidden using the Show Mask button.

The Mask array can be moved as a group by clicking on the Up, Down, Left or Right buttons. An individual box within the array can also be moved by first clicking on that box to select it, then using the Up, Down, Left or Right buttons to move it.

The Mask can be optimally positioned by clicking Optimize Mask. It can then be saved as the mask-data.xml file in the Mask folder of the iQ5 Folder by clicking Save Mask. The Mask can be restored to the last saved Mask by clicking Reload Mask. The default, factory Mask can be restored by clicking Restore Factory Mask.

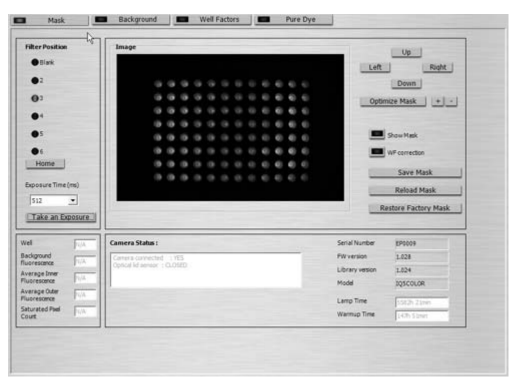


Fig. 6.5. The Mask window.

The main use of the Mask screen is to adjust the masks although there are other uses including capturing an image of the experimental plate to check the response of a probe or to assess the completion of a reaction. With a plate image you can obtain fluorescence readings for each of the 96 individual wells.

6.2.1 Calibration (Alignment) of the Mask

What You Will Need

You need the following to calibrate the iQ5 Camera:

- 1. iCycler iQ External Well Factor Solution (found in the iQ Calibrator Dye Solution Set)
- 2. 96-well PCR plate or preferred reaction vessel (referred to as 'plate')
- 3. Optical quality sealing tape or preferred sealing method

To prepare an external well factor plate:

- 1. Dilute the 10x External Well Factor Solution to 1x with water.
- 2. In the 'plate' pipet equal volumes of the 1x external well factor solution into ALL 96 wells using the volume that you will be using in your experiment.
- 3. Seal the plate using the sealing method you will use in your experiment.

To align the mask:

- 1. Click Home.
- 2. Click the Filter Position 2 option button.
- 3. Click Take an Exposure.
- 4. Click Show Mask.
- 5. If any pink pixels are present in the mask, reduce the exposure time and retake an exposure. Keep retaking the exposure until no pink pixels are present.
- 6. Click Optimize Mask.
- 7. Click Save Mask.

6.3 Background Calibration

Background calibration is performed to account for fluorescence in the experimental system that is due to the reaction vessel and sealing mechanism. Open the Background window by clicking Background as shown below.

instructions:				
Refer to Help File for instructions on how to run this type of celbration.		Seal Type	Film	
	R	Vessel Type:	Plates •	
	-			
		Collect	Background	

Fig. 6.6. The Background window.

What You Will Need

You need the following to calibrate the iQ5 Camera:

- 1. 96-well PCR plate or preferred reaction vessel (referred to as 'plate')
- 2. Optical quality sealing tape or preferred sealing method

6.3.1 Preparation of a Background Calibration Plate

Prepare a background calibration plate by sealing the plate using the sealing method you will use in your experiment.

To perform background calibration:

- 1. Select a well seal.
- 2. Select a vessel type.
- 3. Click Collect Background Factors.

- 4. When the iQ5 software completes the Background Calibration run, a dialog box appears with the message, "Background Calibration Run Complete."
- 5. Click OK to exit.

6.4 Well Factors

Open the Well Factors window by clicking Well Factors as shown below. The Well Factors window is used to collect Persistent Well Factors

instructions: lefer to Help File for instructions on how to run this type of calibration.			Film	-
	40	Seal Type: Vessel Type:	Plates	•
		Sample Volume:	25 -	the second se
		Collect D	ersistent Well Fac	tors 1

Fig. 6.7. The Well Factor Selection window.

Well factors are used to compensate for any system or pipetting non-uniformity in order to optimize fluorescent data quality and analysis. Well factors may be collected directly from the experimental plate (dynamic well factors) or from an external well factor plate (persistent well factors).

The better source of well factors, in terms of correcting non-uniformity, is the actual experimental plate. However, in order to collect well factors from the experimental plate, the plate must meet certain requirements and must be cycled for approximately five minutes extra.

6.4.1 Dynamic Well Factors

In order to collect Dynamic Well Factors on of the following must be true

- 1. All wells must have the same combination of fluorophores AND the same concentration of fluorophores.
- 2. No fluorophore is used both alone and in combination with other fluorphores AND no fluorphore is used in more than one combination of fluorophores.

The collection of dynamic well factors is a completely automated process which begins as soon as the Begin Run button is clicked and the file name saved. When collecting dynamic well factors, the plate is held at 95°C for two and a half minutes prior to the first cycle with a setpoint of 90.0°C or higher. You may wish to adjust the dwell time/temperature for your first step of your thermal protocol accordingly.

6.4.2 Persistent Well Factors

Persistent well factors are not collected from the experimental plate but on a separate calibration plate containing the volume of 1 x External Well Factor Solution in all 96 wells.

In addition the persistent well factors must be collected with the same reaction vessel and sealing mechanism as the experiment will be run with. Persistent Well Factors are store in the Well Factors folder in the iQ5 Folder.

Persistent Well factor files can be viewed by selecting Calibration Data from the View menu.

Files will be named

Persistent_vessel_seal_volume_iQ5CameraSerial#_iCyclerSerial#

where

vessel is one of plates, (tube)strips or tubes

seal is one of film, flat caps or domed caps.

The plate setup used to run the experiment will have a reaction vessel, seal and volume associated with it. This plate setup will not be able to be run in an experiment unless it has the appropriate calibration files including a Persistent Well Factor calibration file.

In general persistent well factors can be used for about one month, but should be collected anytime something pertinent to the optical system is changed such as the optical filters or the camera lamp. A warning will occur when Persistent well factors are older than 30 days.

Persistent Well Factors may be used for any experimental run but there are situations in which they must be used.

6.5 Real-Time PCR Experiments using DNA Binding Dyes

In most real-time PCR experiments using DNA-bining dyes, like SYBR[®] Green I or ethidium bromide, dynamic well factors may not be collected. When the template DNA is denatured, the fluorescence of these dyes is not sufficiently high to calculate statistically valid well factors using the experimental plate.

There are three possible solutions to this problem:

- 1. use iQ[™]SYBR Green Supermix which already includes fluorescein
- 2. use Persistent well factors
- for experiments with SYBR Green I, spike the master mix with fluorescein (the addition of fluorescein provides sufficient fluorescence at 95°C for the collection of well factors from the experimental plate while not interfering with the PCR reaction.

6.5.1 Spiking Real-Time PCR Experiments using DNA binding Dyes with Fluorescein

The Bio-Rad iQ SYBR Green Supermix is already spiked with a small amount of fluorescein that permits the collection of well factors from the experimental plate. It is also possible to collect well factors from the experimental plate with other commercial SYBR Green mixes or with home-brew mixes by adding sufficient fluorescein to bring the reaction mixture to 10 nM fluorescein.

First make a 1 mM solution by a 1:1000 dilution of the 1 mM stock Fluorescein Calibration Dye in PCR buffer (10 mM Tris, pH 8.0, 50 mM KCl, 3 mM MgCl₂). Then add 1 part of the 1 mM fluorescein with 990 ml of master mis to yield a final concentration of 10 nM fluorescein. Once well factors are collected from the experimental plate, they are written to the opd file and the software continues to executer the programmed protocol.

6.6 Pure Dye Calibration - iQ5 Systems Only

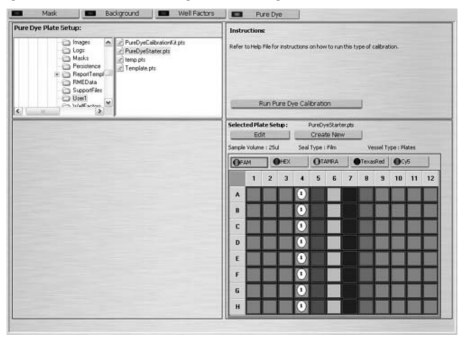


Fig. 6.8. The Pure Dye Calibration window.

Before you perform a Pure Dye Calibration run, complete the following steps:

1. Ensure that the mask has been aligned.

- 2. Ensure that Background Calibration has been performed
- 3. Ensure that Persistent Well Factors have been generated.
- 4. Place the Pure Dye Calibration Plate in the iCycler.
- 5. Select the Pure Dye tab in the Calibration module.

What You Will Need

You need the following to calibrate the iQ5 Camera:

- 1. iCycler iQ Calibrator Dye Solution Set
- 2. iCycler iQ External Well Factor Solution (found in the iQ Calibrator Dye Solution Set)
- 3. 3 x 96-well PCR plate or preferred reaction vessel
- 4. Optical quality sealing tape or preferred sealing method

6.6.1 Preparation of the Pure Dye Calibration Plate

To prepare a pure dye calibration plate:

- 1. Pipet equal volumes of the appropriate 1x calibrator solution into 8 wells of the 96-well plate, using the volumes that you will use in your experiment.
- 2. Repeat for all fluorophores needed.
- 3. Seal the plate using the sealing method you will use in your experiment.

6.6.2 Steps Required for Pure Dye Calibration

To perform a Pure Dye Calibration run:

- 1. Select or Create a pure dye plate setup.
- 2. Click Run Pure Dye Calibration.
- 3. When iQ5 completes the Pure Dye Calibration run, a dialog box appears with the message, "Pure Dye Calibration Run Complete".
- 4. Click OK to exit.

6.6.3 Selecting a Pure Dye Plate Setup

Select a Pure Dye Plate Setup from the file selector window within Pure Dye Calibration tab. When loaded, the fluors on the plate will display within the plate display in the lower right-hand corner of the Pure Dye Calibration tab.

Select	ed Pla	te Sel	tup:										
	Ed	it			Creat	e Nev	v						
Sample	Volum	e : 25u	I	Seal	Type :	Film		Ve	Vessel Type : Plates				
OF/	AM		HEX		OTA	MRA		Texas	TexasRed OCy5				
	1	2	3	4	5	6	7	8	9	10	11	12	
A				1									
В													
С													
D													
Ε				1									
F				1									
G				1									
н				1									

Fig. 6.9. Pure Dye Calibration Plate Setup.

Check the Sample Volume, Seal Type and Vessel Type fields. f these fields match the intended procedure and the correct fluors are displayed, proceed with Pure Dye Calibration. If any of these particulars vary from the intended experiment parameters, the Pure Dye Plate Setup must be edited or a new Pure Dye Plate Setup created.

6.6.4 Editing or Creating a Pure Dye Plate Setup

To edit a Pure Dye Plate Setup, select the Pure Dye Plate Setup that most closely resembles the experimental parameters desired and click the Edit button. The Pure Dye Plate Setup Editor will load:

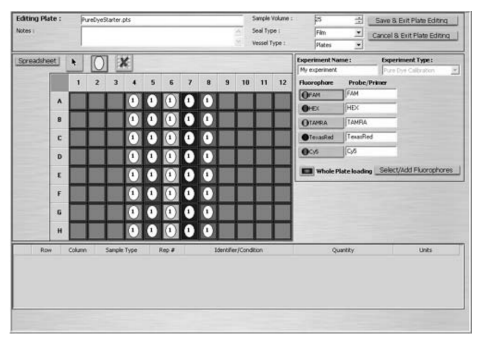


Fig. 6.10. Creating a Pure Dye Calibration Plate Setup.

Select/Add Fluorophores

To choose the fluors needed to calibrate the iQ5 instrument, select from the predefined fluorophores or add custom fluors by clicking the Select/Add Fluorophores button. The Fluor Selector will display.

	Filter Position	Fluor	Selected	Color
	2	FAM	7	Line
		SYBR	Г	
		SY8R1		
		SYBR2	Г	
		SYBR3	Г	
		SYBR4	Г	
		SYBR5	Г	
	3	HEX	1	Green
		JOE	Г	1177.0
		TET	Γ.	
		VIC	E .	
	4	Cy3	Г	
		TAMRA	1	Skyfikar
1	5	TexasRed		Blue
•				
	-			
Fluor	Name :			-
Filter	Position:	2	*	Cancel

Fig. 6.11. The Select/Add Fluorophores Dialog Box.

Add a fluor from the list displayed by clicking the check box next to the fluor name. To add a custom fluor (not in the list already), first specify the filter pair which will analyze them from the drop-down menu. The choices are

- Position 2 485/20X 530/30M Fluorescein (FAM),SYBR Green I
- Position 3 530/30X 575/20M HEX, TET, VIC, JOE
- Position 4 545/30X 585/20M TAMRA/Cy3
- Position 5 575/30X 625/30M Texas Red, ROX
- Position 6 630/30X 685/30M Cy5, LC640

Then, specify the name and color of the custom fluor by typing the name in the Fluor Name field and clicking Add New Fluor. The new fluor name appears in the list for the selected filter pair. Click the check box next to the new fluor name, then select the color to specify the fluor in the graphic by clicking on the color field that appears. Choose contrasting colors for maximum clarity, then Click OK.

Note: Two or more different dyes may be used with the same filter pair; however, two dyes using the same filter pair may not be used in the same well.

Select new Sample Volume, Seal Type and Vessel Type from the drop-down menus as appropriate for this experiment, then click Save & Exit Plate Editing. Choose a meaningful file name, and save the Pure Dye Plate Setup.

Important: Pure Dye Calibrations are made for each machine, dye type, sample volume, seal type and vessel type, and are machine- and experiment-specific. Do not attempt to use Pure Dye Calibration results from one machine on a different machine, or different seal, vessel and sample volume pure dye calibration results on the same machine.

6.7 Viewing Calibration Files

View	Reports	Tools
Log Cal	js ibration Da	ata

Calibration files can be viewed by selecting Calibration Data from the View menu. Background factors file will be found in the Background Factors folder in the iQ5 Program Folder. Persistent Well Factors will be found in the Well Factors folder in the iQ5 Program Folder. Pure Dye Calibrations will be found in the RMEData folder in the iQ5 Program Folder.

Section 7 User Profiles

To assist with the management of files and data created by the iQ5 software, one or more User Profiles may be created for users of the iQ5 or MyiQ system.

User Profiles in the iQ5 software consist of the following information:

- Login name required
- Password optional
- Permission settings for collecting, saving, and analyzing data
- Preferred defaults (User Preferences) for collecting, saving, and analyzing data

Note: All User Profile information resides on the hard drive of the computer where the software has been installed, this means that User Profile information can NOT be shared over a network.

When the software is installed, the default User Profile has the ability to add and configure additional users through the User Profile module. The User Profile module is where users can be added, their permissions controlled and their preferences set. This module of the iQ5 software consists of 2 tabs:

- User Preferences
- Administration

7.1 User Preferences

User Preferences, found in the User Profile module, can be used to customize the preferences of the currently logged in User.

For each User created in the iQ5 software, user-specific preferences can be set for:

- File Paths
- Plate Setup options
- Protocol options
- File Selection at application startup
- PCR Quant module analysis settings
- Allelic Discrimination module analysis settings
- End Point module analysis settings
- Gene Expression module analysis settings

7.1.1 File Paths Preferences

Each file created by the iQ5 software is destined to a specific folder location on your computer, this is known as a "file path". The File Paths preferences box can be used to select the folders that you wish to save your files into when logged into the iQ5 software. The default File Paths for all users is a folder created with the current user's User Name, for example:

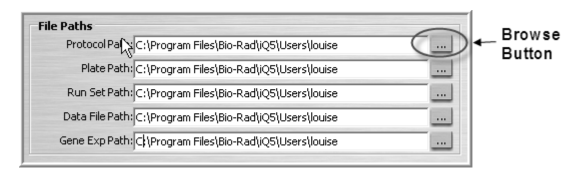


Fig 7.1 The File Paths dialog box.

File Paths can be specified for:

- Protocol files
- Plate Setup files
- Run Sets
- Data Files
- Multifile Gene Expression Analysis Files (Gene Exp Path)

To set the file path to an existing directory, click the browse button and navigate to the appropriate folder.

To set the file path to a new directory, click the browse button, then click Make New Folder in the Browse For Folder dialog box. A new folder will appear within the currently selected folder. To rename this folder, right-click, select Rename, and then enter the new name of the folder in the text field next to the newly created folder. Click OK when finished.

Browse For Folder	?×
Persistence	
E Constante E Constante Report Templates	
🔂 RMEData	
🛅 SampleFiles	
🛅 SupportFiles	
😑 🧰 Users	
🚞 admin	
New Folder	~
	>
Make New Folder OK Car	icel

Fig 7.2 Creating new folders in the Browse For Folder dialog box.

Note: Changes to default File Paths DO NOT become active until the software has been restarted.

7.1.2 Plate Setup Preferences

The Plate Setup preferences box can be used to define the default conditions when a User creates a new plate setup.

Plate Setup		
Concentration Units: copy number	✓ Seal Type: Film	
Vessel: Plates	▼ Sample Volume: 25 ÷	
Scientific I	Notation	
Dilution Sctor: 10 📑		
Dilution Series: Increasing 	Decreasing	
Replicate Size: 1	Replicate Loading: Row & Column]
Calibration Plate: D:\Program Files	s\Bio-Rad\iQ5\Users\louise\puredyestarte	
		-
Fluorophores: Cy3	LC640 TET	
✓ Cy5	✓ TexasRed VIC	
✓ FAM	ROX SYBR1	
✓ HEX	SYBR SYBR2	
JOE	TAMRA SYBR3	
<		

Fig 7.3. Definition of Plate Setup preferences

The following default conditions can be set for the Plate Setup:

- Concentration Units: Choose from 14 different units of measure to match the units used in your experiment.
- Seal Type: Chose from Film, Domed Cap and Flat Cap.
- **Vessel Type**: Chose from Plates, Strips (tube strips) and Tubes.
- Sample Volume: Set the sample volume (in microliters) that will be used as the default value.
- **Dilution Series Settings**: Set the defaults for the Dilution Factor, and specify whether the series is Increasing or Decreasing in quantity. You may specify if Scientific Notation is to be used in the display of the Dilution Series values.
- Replicate Size: Used to specify the default replicate size.
- **Replicate Loading**: Used to specify the default direction replicates will be loaded. You can choose from Row (replicates added horizontally), Column (replicates added vertically), and Row and Column (replicates added in a block). See Specifying Replicates for more information.
- **Calibration Plate**: This is used to set the default file to be selected for the Pure Dye Calibration plate. This file will be used after Application Startup.
- **Fluorophores**: Place a checkmark next to the fluorophores that you wish to have automatically selected when creating a new plate setup. No more than 5 fluorophores can be selected. At least one fluorophore must always be selected.

7.1.3 Protocol Preferences

The Protocol Preferences box can be used to set the defaults for the current User for Melt Curve, Gradient and Well Factor Collection Type.

÷	End Temp: 95.0	+	Temp Change: 0.5	÷
÷	Range: 10.0	<u>+</u>		
Collection T	ype: Persiste	ent	Opnamic	
	÷	Range: 10.0	- Range: 10.0 -	Range: 10.0

Fig 7.4. Defining user specific Protocol options.

The following default conditions can be set for the Protocol

- **Melt Curve**: Enter the desired beginning temperature (Set Point), the Ending Temperature (End Temp) and the change in temperature that will occur at each repeat (Temp Change). Click Save.
- **Gradient**: Enter the desired lowest temperature (Set Point) and the Range for the gradient in the appropriate text boxes (or use the scroll buttons to enter these values). Click Save.
- Well Faction Collection Type: Select the radio button associated with the Well Factor that you wish to use as the default type. Click Save. If Dynamic well factors are chosen as the default type they will be used unless restricted by the Plate Setup.

7.1.4 File Selection at Application Startup Preferences

The file that you wish to have automatically selected when the iQ5 software is opened is determined using the File Selection at Application Startup radio buttons.



Fig 7.5. Selecting custom file defaults for application startup.

- Last File Used: This option will select the last Protocol, Plate Setup, Run Set or Data File used as the file to display in the Workshop screen when the iQ5 software starts up.
- **Data File**: If you wish to have a specific data file selected when the iQ5 software opens, select the Data File radio button and then use the Browse button to navigate the data file that you wish to have selected on application startup.
- **Run Set**: If you wish to have a specific Run Set file selected when the iQ5 software opens, select the Run Set radio button and then use the Browse button to navigate the run set file that you wish to have selected on application startup.
- **Protocol/Plate**: If you wish to have a specific Protocol or Plate Setup file selected when the iQ5 software opens, select the Protocol/Plate radio button and then use the Browse button to navigate either the Protocol or Plate file that you wish to have selected on application startup.

7.1.5 PCR Quant Screen Preferences

The PCR Quantification preference box can be used to set the defaults for the PCR Quant screen. You can set the defaults for four major settings that will impact the analysis or display of data collected from your real time PCR detection system.

PCR Quant Baselining:	Auto Calculated	OUser Defined	
	ned Baseline Cycle Range;	to 10	-
Log View:	●On ●Off		
Analysis Mode:	PCR Base Line Subtracted (Curve Fit 💽	
Grid Rows: 💻 I	dentifier 💻 Ct 📕	Concentration	EP Call

Fig 7.6. Defining user specific PCR Quant analysis and display options.

- **Baselining**: By default, PCR baselines are auto-calculated. With the Baselining User Preference setting, you can override auto-calculation by clicking User Defined. After you click User Defined, the iQ5 software activates the User Defined Baseline Cycle Range boxes. Enter the desired values for the start cycle and ending cycle for all traces. Auto Calculated is the factory default.
- Log View: Use this option to select either a semi-logarithmic or linear view of the PCR Amplification Chart data. With the Log View option On, a semi-logarithmic view of the Amplification data will be displayed. Log View: Off is the factory default.
- **Analysis Mode**: You can select from three analysis modes in the Analysis Mode drop-down list boxs. The three analysis modes include: Background Subtracted, PCR Base Line Subtracted, PCR Base Line Subtracted Curve Fit. PCR Base Line Subtracted Curve Fit is the factory default.
- **Grid Rows**: The Grid Rows options are a set of check boxes that you can use to display additional details about the data displayed in the PCR Quant screen. These optional details will be display in the results spreadsheet of the PCR Quant screen, and include the following information about each sample: Identifier, Concentration, Threshold Cycle, and End Point Call.

7.1.6 Allelic Discrimination Module Preferences

The Allelic Discrimination preference box can be used to set the default display conditions used in the Allelic Discrimination screen. The Display Mode can be set to Threshold Cycle or RFU using the radio buttons. The radio buttons for the Normalize Data option only become active when the Display Mode is set to RFU.

Allelic Discrimination			
Display Mode: Threshold Cycle	●RFU	Normalize Data: 🗨 Yes	●No

Fig 7.7. Defining Allelic Discrimination display options.

7.1.7 End Point Module Preferences

The End Point module preference box allows you to predetermine several analysis and display options associated with analyzing final RFU in an End Point analysis assay.

Method: Negativ	ves	•	Ranks: 10	+
End Cycles to Avg:	PCR:5	÷	End Point Only Run: 2	+
Tolerance Mode: RFUs		•	Tolerance %: 10.0	÷
Color Scheme: Yellow	Green	•		

Fig 7.8. Defining analysis and display options for the End Point module.

- **Method**: The Method box allows you to select the method of assigning positive and negative values to your unknowns based on RFU values. The Method box consists of the following three choices: Negatives, Positives, and Positives & Negatives. Analysis by comparison to negative controls (Negatives) is the default method.
 - **Ranks**: The number of Ranks allows assignment of samples into distinct groups based on their RFU values. The default Rank value is 10 and the minimum number of Ranks is 3.
- End Cycles to Avg: End Cycles to Average is the number of cycles, from the last cycle, that will be used to calculate an average End Point RFU value. The end cycles to average field defaults to 5 for non-End Point only (PCR) runs.
 - End Point Only Run: The end cycles to average field defaults to 2 for End Point only runs.
- Tolerance Mode: End Point Tolerance defines the margins for sorting unknowns as positives or negatives. The End Point Tolerance drop-down list box consists of two choices: RFUs, and Percent of Range.
 - **Tolerance %**: Select a value between 1 and 99 percent for this setting. The default percent of range tolerance is 10 percent.
- Color Scheme: Colored rank boxes, symbolize the number and order of ranks in the End Point Analysis. There are five color-options to choose from that allow a change in the color scheme of the rank boxes, once the data are analyzed.

7.1.8 Gene Expression Module Preferences

The Gene Expression module preference box allows you to predetermine several analysis and display options associated with analyzing Ct values for Gene Expression analysis.

elative to:	● Zero	Control	
	• Condition	Gene	
YAxis:	●Log 2	OLinear	
Scaling:	Highest	 Lowest 	• Unscaled
Method:	• ddCt	●dCt	Std Devs: 1.00 ÷

Fig 7.9. Defining analysis and display options for the Gene Expression module.

- **Relative to**: These options, allow you to present data with axes originating at 1 (relative to control) or at zero (relative to zero).
- X Axis: This option allows you to graph either genes or conditions on the x-axis.
- **Y** Axis: This option allows you to display the graph with the y-axis in a log2 scale or on a linear scale.
- Scaling: "Scale to control" is another scaling option which is accomplished by assigning a control in the Condition List.
- **Method**: These options allow you to set the default analysis mode for your Gene Expression analysis. You can select Normalized Expression (Ct) or Relative Expression (Ct) as a default analysis setting.
- **Std Devs**: The default presentation for the error bars is plus and minus one standard deviation. You can change the multiplier to get plus and minus 2 or 3 standard deviations.

7.2 User Administration

The Administration tab can be by a user with an "iQ5 Administrator" Role to:

- Add or delete users
- Edit user details (i.e., Full Name and eMail information)
- Set or change passwords
- Control the permissions of each User Role

Only an iQ5 Administrator can access and modify information contained in the Administration window.

Users specified as iQ5 Principal, iQ5 Operator, and iQ5 Guest have read-only access to the Administration window, where they can review permission settings for their role.

7.2.1 Adding New Users

Users are added using the Defining Users spreadsheet.

Defining Users:			Save User Changes			
User Name	Full Name	Role		eMail	Password	Delete
admin	Administrator	iQ5 Administrator	~		******	
clopez		iQ5 Principal	*		*****	
ksmith		iQ5 Operator	~			
Russell Lab		iQ5 Principal	~			
Summer Students		iQ5 Guest	~			
			~			



- User Name: REQUIRED The User Name is a set of alphanumeric characters that uniquely defines each user. It can be up to 15 characters long and composed of upper and/or lower case characters.
- Full Name: OPTIONAL Use this field to specify the full name corresponding to the User Name
- **Role**: REQUIRED There are 4 Roles in the iQ5 Software: Administrator, Operator, User and Guest. Each of these Roles gives users within that role permission to access specific features and functions of the software. Permissions granted to all of the Roles, with the exception of Administrator, can be customized by the Administrator
- **eMail**: OPTIONAL The eMail cell is an informational area for contact information.
- **Password**: OPTIONAL The password can be any combination of letters, numbers or symbols. It can be of any length. An Administrator can also change the password of Users. This is useful if a User forgets their password and a new password must be assigned.

To ADD a new user:

- 1. Type the desired login name into an empty User Name box of the Defining Users spreadsheet.
- 2. Optional: Type in the Full Name of the User.
- 3. Define the Role of the User by using the pull-down menu in the Role cell. A User can be an Administrator, Operator, User or Guest. The features/functions that each of these User Roles is permitted/ restricted to use is defined by a user with an Administrator Role using the Defining Roles spreadsheet.
- 4. Optional: Type in email or phone contact information of the User.
- 5. Optional: Type in a password for the user. This password, defined here by the Administrator, can later be changed by the User using the Change Password option found in the Tools menu.
- 6. Click Save User Changes.

Adding a user will create a folder with the new user's name. This folder will be created in the Bio-Rad/iQ5/Users folder and will be the default folder for all files saved when this User is logged into the software. The user may as unless changed in User Preferences.

7.2.2 Deleting Existing Users

Users are removed using the Defining Users spreadsheet.

- 1. In the Defining Users spreadsheet, check the box in the Delete column for the user that is to be removed.
- 2. Click Save User Changes.

Note: No messages will appear to confirm this action, the User Profile will immediately be removed from the Defining Users spreadsheet.

7.3 Logging on to the iQ5 Software

There are two ways for a User to sign in to the iQ5 software:

- 1. At the start up of the iQ5 program.
- 2. Through the Switching Users option, found in the Tools menu.

🕅 Login		×
R		
User:	Russell Lab	
Password:		
		<u>O</u> K <u>C</u> ancel

Fig 7.11. The User Login dialog box.

The User Name can be selected from the pull-down menu in the Login dialog box. Enter the password and click OK to log on to the iQ5 Software.

If the User Name or Password in incorrect then the following message will be displayed:

Login fa	Login failed					
\otimes	Make sure you have the correct password and user name, taking care to type the password correctly including capitalization. Make sure the Caps Lock has not been accidentally turned on.					

Fig 7.12. Incorrect login error message.

Click OK and enter the correct information. If you have forgotten your password, it may be reset by a user configured as an iQ5 Administrator.

The current user can be determined by looking title bar (in the top-right corner) of the software. The current user will be displayed in parentheses next to the Bio-Rad iQ5 header:

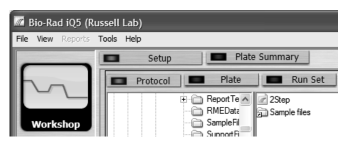


Fig 7.13. User status display in iQ5 title bar.

7.3.1 Switching Users

To switch from the currently logged in User click on the Workshop module icon, then select Switch User from the Tools menu.

The Login dialog box will appear (see Fig 7.11)

Select the desired User from the pull-down list, enter the password (if any) in the Password text box and click OK.

If the User name you want is not available you will need to add it in the User Profile module, Administration tab.

7.3.2 Changing Password

The password of the current user can be changed by selecting the Tools menu, Change Password selection.

Enter your Old Password in the top text box, enter your new password in the New Password and Confirm New Password text boxes. Click OK

Gri
J
OK Cancel

Fig 7.14. The Change Password dialog box.

7.3.3 Defining Roles

Defining Roles:	Save Role Changes		
Permission	iQ5 Principal	iQ5Operator	iQ5 Guest
Save Any File	Image: A start of the start	1	×
Start, Pause and Abort Runs	 Image: A start of the start of	4	>
Add Repeats to a Run	¥	1	
Perform Skip Cycles	\checkmark		
Create Gene Study Files	¥		×
Perform Instrument Calibration	\checkmark	1	
Apply Different Calibrations to a Data File	¥	1	
Change Endpoint Run Set Point	\checkmark	1	
Use Expired Calibrations for Runs	¥	1	

Fig 7.15. The Defining Roles interface of the User Profile module.

Listed above is the set of features and functions (permission) that can be granted by the iQ5 Administrator to users assigned to one of three non-Administrator Roles. The Administrator can customize the Roles of a Principal, Operator and Guest by checking or unchecking the box associated with each function and each Role. To apply the selected changes click the Save Role Changes button, to restore factory default roles click Restore Factory Roles.

Note: In the factory default roles, users assigned to the iQ5 Guest Role do not have any permissions granted to them. By default, these users are granted "read-only" access to the software unless the ability to start runs and save files is granted by the iQ5 Administrator.

Viewing User Permissions

To view the current permissions settings for existing iQ5 Users, access the User Administration window by clicking on the Administration button.

All non-Administrator users will have read-only access to the information in the User Administration window

Section 8 Maintenance of the iQ5 and MyiQ Real Time PCR Detection Systems

8.1 Cleaning the iQ5 and MyiQ Real-Time PCR Detection Systems

Take care not to spill liquids onto or into the iCycler thermal cycler or the optical module.

Cleaning may be done using a lint-free cloth or paper towel. The case may be cleaned using a soft, lint-free cloth and water.

8.2 Filter Description and Installation Instructions

The filters designed for use in the iQ5 and MyiQ optics modules are made of glass and mounted in plastic holders. The filter holders are held in either the excitation or emission filter wheel of the optics module. Each filter wheel holds six filters. Every position in a filter wheel must have a filter or an opaque filter blank to avoid damage to the CCD detector. The first position in each filter wheel is designated as the "home" position and must always contain an opaque filter blank. Filters can be removed for cleaning or replacement. If a filter shatters or breaks during the installation process, contact your local Bio-Rad office immediately for service, do not attempt to remove the broken components from the interior of the camera housing.

It is critically important that the excitation and emission filters are in the correct positions in the filter wheels, please confirm that the filters are in the proper location after cleaning or replacing filters in the optics module. The table below summarizes the positions of the filter pairs, their optical characteristics, and the recommended fluorophores with which they are compatible.

8.2.1 Filter Positions and Recommended Fluorophores

- 2 485/20X 530/30M Fluorescein (FAM),SYBR Green I
- 3 530/30X 575/20M HEX, TET, VIC, JOE
- 4 545/30X 585/20M TAMRA/Cy3
- 5 575/30X 625/30M Texas Red, ROX
- 6 630/30X 685/30M Cy5, LC640

Note: The filter designation 485/20X indicates that this filter will allow light between 475 and 595 nm to pass through. The first number, 485, indicates the center of the wavelength of light. The second number, 20 indicates the total breadth of wavelengths of light that can pass through it. The letter "X" indicates that the filter is specified for excitation only, and the letter "M" indicates emission only types of filters. Excitation and emission filters are not interchangeable.

8.2.2 Accessing the Existing Filters

To access the existing filters, proceed as follows:

- 1. Turn off the power to the optics module.
- 2. Reach behind the optics module and unscrew the short fastener that secures the lid of the module in place.
- 3. Using gentle pressure and both hands, push inward on the rear vents located on the top half of the casing. Lift upward to remove the cover of the optics module.
- 4. To access the excitation filter wheel, remove the black plug from the slot located near the lamp, at the right-hand side of the optics module. To access the emission filter wheel, remove the plug from the slot located at the top of the instrument.
- 5. Turn the filter wheels to the desired positions using the supplied ball-end hex driver. As long as the power to the optics module is off, the filter wheels may be turned freely in either direction.
- 6. To remove a filter, grasp it on both sides with the filter removal pliers and squeeze the tab in; gently pull the filter up and out.
- 7. To insert a filter, grasp the filter with the pliers and insert it into a vacant slot. For the excitation filters, the tab on the filter should face toward the front of the instrument. For the emission filters, the tab on the filter should face the right of the instrument. Be sure that every position in the filter wheel has either an excitation or emission filter or a filter blank before powering on the system.
- 8. After the filters or filter blanks have been inserted, replace the rubber plugs over the slots of the filter wheels.
- 9. Realign the tabs on the front end of the optics module cover with the tabs on the main housing. Lower the cover until the top half of the camera housing snaps into place.
- 10. Replace the screw in the rear of the optics module to secure the casing.

8.2.3 Cleaning the Filters

The iQ5 and MyiQ instruments are shipped with the specified filters pre-installed, ready for use. Normally, you should not have to reconfigure or replace the optical filters that come with your unit. However, the optical filters are removable and user-serviceable by design.

Handle the optical filters with care as they can crack if dropped. Also, avoid touching the surfaces of the filters, especially with fingers, as this can impair data quality. Contact Bio-Rad for replacement filters.

You may clean the optical filters by wiping them gently with lens paper and 70% ethanol. You may obtain lens paper and lens cleaning solution by ordering a Lens Cleaning Kit from Bio-Rad (catalog #170-7731). After cleaning both sides of an optical filter, hold it up to the light and make sure that no smudges, fingerprints, debris, or water marks remain. Once clean, you may reinsert the optical filters as described above.

8.3 Replacing the Lamp

When replacing the lamp, you must only use lamps supplied by Bio-Rad. Bio-Rad lamps are subject to additional tests and standards geared specifically toward Real-Time data collection. Lamps from alternative sources, which may appear to be similar, may not deliver the same optical quality, performance, and lifetime as those supplied by Bio-Rad. If the camera is to remain continuously on, we recommend the lamp to be replaced every 6 months.

Note: When a lamp is overdue for replacement, if may flicker sporadically, causing a wavering of the data in all wells simultaneously. Installing a new lamp will alleviate this problem.

Caution: Take care when changing the lamp as it may be hot. Allow at least 15 minutes after turning off the optics module before removing the lamp.

The lamp is located on the right side of the optical module.

8.3.1 Lamp Replacement Procedure

- 1. Turn off the power to the optical module.
- 2. Reach behind the optical module and unscrew the short fastener that secures the lid of the module in place.
- 3. Using a gentle pressure with both hands, push inward on the rear vents located on the top half of the iQ5 casing. Lift upward to remove the cover of the optical module.
- 4. Push up on the lamp spring clip to release the lamp from the bracket.
- 5. Lift the lamp out of the socket.
- 6. Install the new lamp using the reverse of steps 1-5. Hold the new lamp by the outer reflector and do not touch the bulb. Be sure the spring clip is down before inserting the lamp into the socket. Push the lamp firmly into the bracket, then close the case and secure the lid.

Appendix A Support

The iQ5 real-time PCR detection system, and MyiQ real time PCR detection system are warranted against defects in materials and workmanship. For specific warranty information, contact your local Bio-Rad office.

If any defects should occur during the warranty period, Bio-Rad will replace the defective parts without charge. However, damage or defects resulting from any of the following causes are specifically excluded:

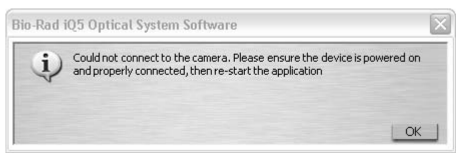
- 1. Improper operation.
- 2. Use of improper solvent or sample.
- 3. Use with tubes, plates, or sealing materials not specified by Bio-Rad Laboratories for use with the iQ5 or MyiQ real-time PCR detection system.
- 4. Deliberate or accidental misuse.
- 5. Repair or modifications done by anyone other than Bio-Rad Laboratories.
- 6. Natural disaster of any kind.

The warranty does not apply to fuses.

For inquiry or request for repair service, contact Bio-Rad Laboratories after confirming the model and serial number of your instrument. For technical support, call your local Bio-Rad office, or in the US, call 1-800-4BIORAD (1-800-424-6723), or visit us on the Web at discover.bio-rad.com.

Appendix B Error Messages

B.1 Software Startup

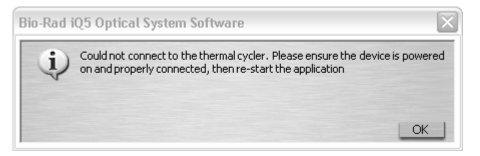


Cause: The software is unable to communicate with the camera.

Solution: Ensure that the camera is on.

Check that the USB cable is securely connected to both the camera and computer.

Ensure that the USB port in the computer is 2.0 High Speed. You may need to close the software and restart both the camera and iCycler base before reopening the software.



Cause: The software is unable to communicate with the iCycler base.

Solution: Ensure that the iCycler base is turned on.

Check the serial cable connection between the iCycler base and the Camera. You may need to close the software and restart both the camera and iCycler base before reopening the software.

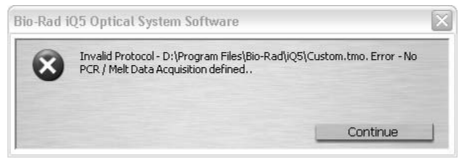
Bio-Rad io	Q5 Optical System Software
?	Improper shutdown detected, if a run was in progress during the last session, it could be still in progress, starting the application will abort any run. Continue?
	Yes No

Cause: The iQ5 Software did not shutdown properly.

Solution: If the software had been controlling a run check the iCycler base to see if the run is still proceeding. If a run is still proceeding clicking Yes to this dialog box will abort the run, clicking No will allow the run to continue but will not re-start the software.

If the software is not connected to a camera, for example on a computer which is used only to analyze the data, clicking Yes will re-start the software.

B.2 Protocol and Plate Errors



Cause: There is no step defined for data collection or analysis in the thermal protocol. At least one data

acquisition step must be present in the thermal protocol.

Solution: Specify a data collection at one step of the protocol. A data collection step is shown with a yellow or green camera icon at the appropriate step in the graphical display of the protocol.

Bio-Rad iQ5	Optical Sy	/stem Sof	tware			X
?) ^{Se}	lected fluorop	ohore(s) not	used on plate	e, Remove	unused fluor	rophores?
				1	Yes	No

- Cause: You have selected fluorophores but have not defined any wells to contain these fluorophores
- Solution: This if a reminder to check your plate setup. Clicking No will return you to the Editing Plate window. Clicking Yes will remove the unused fluorophores and save the plate setup.



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 Israel 03 962 3053 200
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