# QuantStudio<sup>™</sup> Design and Analysis Desktop Software USER GUIDE

Getting started with design and analysis of experiments in the desktop software v1.6.

for use with: QuantStudio<sup>™</sup> 1 Real-Time PCR System QuantStudio<sup>™</sup> 3 Real-Time PCR System QuantStudio<sup>™</sup> 5 Real-Time PCR System Publication Number MAN0010408 Revision E.0



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Revision	Date	Description
E.O	30 January 2024	<ul> <li>v1.6</li> <li>Network and password security requirements were added.</li> <li>A chapter was added to describe the use of the software with the Security, Auditing, and E-signature Administrator Console.</li> <li>Additional instructions were provided for exporting run data and results ("Export experiments or results" on page 42).</li> <li>The preferences were updated to include selecting the data destination for different file types ("Set global desktop software preferences" on page 15). This replaces selecting a single default folder or selecting the last location where a file was saved.</li> <li>The instructions to print a report were updated to note that the grouping in the well table is retained in the report ("Print a report of run data and results" on page 44).</li> <li>Instructions to save as were updated to include a new dialog box.</li> <li>The required permissions were included in the applicable sections when security is enabled in the software.</li> </ul>
D.0	22 May 2023	The references to the specific user guides for each real-time PCR instrument were removed.
C.0	19 October 2018	Updated to add QuantStudio™ 1 Real-Time PCR System information.
B.0	17 December 2015	Updates include: <ul> <li>Combination of define and assign functions into a single <b>Plate</b> tab</li> <li>Display of VeriFlex<sup>™</sup> Zones on plate layout</li> <li>Real-time data monitoring in the <b>Run</b> tab</li> <li>Security, Audit, and E-Signature (SAE) features</li> <li>Implementation of locked workflow</li> <li>Selection of an instrument before starting a run</li> <li>Ability to select multiple targets in the results view</li> <li>Various minor changes to the user interface</li> </ul>
A.0	17 April 2015	New document.

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# **Product information**

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# **Product description**

The QuantStudio<sup>™</sup> Design and Analysis Desktop Software allows the user to open, run, and analyze experiments generated with QuantStudio<sup>™</sup> 1 Real-Time PCR System, QuantStudio<sup>™</sup> 3 Real-Time PCR System, and QuantStudio<sup>™</sup> 5 Real-Time PCR System. The software also allows you to set up experiments, send experiments to the instrument, collect data, and analyze the collected data.

The software can be used with the Security, Auditing, and E-signature Administrator Console. The console enables system security by controlling user access, facilitating auditing, and supporting e-signatures. For more information, see Chapter 9, "Use the software with the Security, Auditing, and E-signature (SAE) Administrator Console".

# Network and password security requirements

#### Network configuration and security

The network configuration and security settings of your laboratory or facility (such as firewalls, antivirus software, network passwords) are the sole responsibility of your facility administrator, IT, and security personnel. This product does not provide any network or security configuration files, utilities, or instructions.

If external or network drives are connected to the software, it is the responsibility of your IT personnel to ensure that such drives are configured and secured correctly to prevent data corruption or loss. It is the responsibility of your facility administrator, IT, and security personnel to prevent the use of any unsecured ports (such as USB, Ethernet) and ensure that the system security is maintained.

#### **Password security**

Thermo Fisher Scientific strongly recommends that you maintain unique passwords for all accounts in use on this product. All passwords should be reset upon first sign in to the product. Change passwords according to your organization's password policy.

It is the sole responsibility of your IT personnel to develop and enforce secure use of passwords.



#### **Recommendations for passwords**

Thermo Fisher Scientific recommends enabling a password policy for SAE user accounts with the following minimum number of characters.

- Administrative users-12 characters
- Non-administrative users 8 characters

The use of a password manager is recommended to help create secure passwords.

# Features in the QuantStudio<sup>™</sup> Design and Analysis Desktop Software

Actions	Unlocked template	Locked template <sup>[1,2]</sup>
Properties tab		
Edit experiment name; enter / scan plate barcode; enter user name	1	1
Select instrument / block type, experiment type, chemistry, run mode	1	—
Enter reagent information (chemistry details)	1	1
Method tab		
Edit the thermal protocol, reaction volume, optical filter selection	1	_
Plate tab		
Quick Setup subtab		
Define plate attributes	1	_
Define or assign samples	1	1
Define or assign targets or SNP assays	1	_
Advanced Setup subtab		
Define samples	1	1
Assign samples	1	1
Define targets or SNP assays	1	_
Assign targets or SNP assays	1	<ul> <li>Image: A start of the start of</li></ul>
Run tab		
Start and monitor a run in progress	<ul> <li>✓</li> </ul>	1
View time remaining and plots <sup>[3]</sup>	1	✓



#### (continued)

Actions	Unlocked template	Locked template <sup>[1,2]</sup>
Results tab		
Review run results (analyzed run data)	1	✓
Configure analysis settings	1	—
Export tab	-	
Select export options for run data and run results (analyzed run data)	1	1
Export run data <sup>[3]</sup> ; export template settings	1	✓

<sup>[1]</sup> If you enter the password for a locked template, all listed actions are available.

<sup>[2]</sup> Always save a backup *unlocked* version of a template before saving it as a locked template.

<sup>[3]</sup> This feature is also available from the instrument touchscreen.

# **Experiment types**

Purpose	Description
Standard curve experin	nent
Determines absolute target quantity in	1. The software measures amplification of the target in a standard dilution series and in test samples.
samples.	<b>2.</b> The software generates a standard curve using data from the standard dilution series.
	<b>3.</b> The software uses the standard curve to interpolate the absolute quantity of target in the test samples.



Purpose	Description
Relative standard curve e	xperiment
Determines relative target quantity in samples.	1. The software measures amplification of the target of interest and of an endogenous control target in a standard dilution series, in a reference (calibrator) sample, and in test samples.
	The endogenous control is a target that is expressed equally in all samples; examples of endogenous controls are $\beta$ -actin, GAPDH, and 18S ribosomal RNA. The software can algorithmically incorporate multiple endogenous control targets in relative quantification calculations.
	The reference sample is used as the basis for relative quantification results (or 1× sample). For example, in a study of drug effects on gene expression, an untreated control is an appropriate reference sample.
	2. The software generates standard curves for the target of interest and the endogenous control using data from the corresponding standard dilution series.
	<b>3.</b> The software uses the standard curves to interpolate the quantities of the target of interest and the endogenous control in each sample. The target quantity in each sample is then normalized to the sample's endogenous control quantity.
	<b>4.</b> To determine the relative quantity of the target in test samples, the software divides the normalized target quantity in the sample by the normalized target quantity in the reference sample.
Comparative $C_t$ ( $\Delta\Delta C_t$ ) ex	periment
Determines relative target quantity in samples.	<b>1.</b> The software measures amplification of the target of interest and of an endogenous control target in a reference (calibrator) sample and in test samples.
	The endogenous control is a target that is expressed equally in all samples; examples of endogenous controls are $\beta$ -actin, GAPDH, and 18S ribosomal RNA. The software can algorithmically incorporate multiple endogenous control targets in relative quantification calculations.
	The reference sample is used as the basis for relative quantification results (or 1× sample). For example, in a study of drug effects on gene expression, an untreated control is an appropriate reference sample.
	<b>2.</b> The measurements for the target of interest are normalized to the endogenous control.
	<b>3.</b> To determine the relative quantity of the target in test samples, the software compares the normalized $\Delta C_q$ ( $\Delta C_t$ or $\Delta C_{rt}$ ) for the sample to the normalized $\Delta C_q$ ( $\Delta C_t$ or $\Delta C_{rt}$ ) for the reference sample.

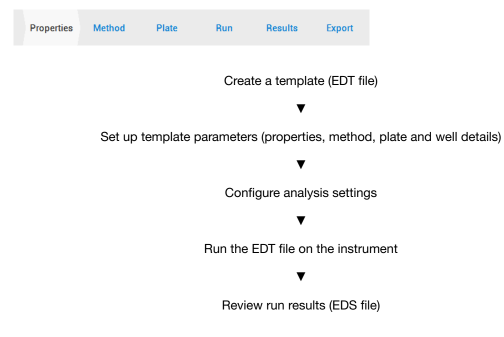
	<u> - 1</u>	$\sim$	-
	k	4	
		16	
		1	
-			

Purpose	Description				
Genotyping experiment					
Detects single nucleotide polymorphism (SNP)	Genotyping experiments use preformulated TaqMan <sup>™</sup> SNP Genotyping Assays that include the following components:				
variants of a target nucleic acid sequence.	<ul> <li>Two sequence-specific primers for amplification of sequences containing the SNP of interest</li> </ul>				
	• Two allele-specific TaqMan <sup>™</sup> probes for Allele 1 and Allele 2				
	1. The software normalizes the fluorescence of the reporter dyes to the fluorescence of the passive reference dye in each well.				
	<ol> <li>The software plots the normalized reporter dye signal of each sample well on an Allelic Discrimination Plot, which contrasts the reporter dye intensities of the allele-specific probes.</li> </ol>				
	<b>3.</b> The software algorithmically clusters the sample data, and assigns a genotype call to the samples of each cluster according to its position on the plot.				
Presence/absence experiment					
Determines the presence or absence of a target nucleic acid sequence in a sample.	The software calls the target present or absent based on an algorithmically determined call threshold. (The call threshold is different from the $C_t$ threshold; the $C_t$ threshold is not used to make calls.)				
Melt curve experiment					
Determines the melting temperature (T <sub>m</sub> ) of the amplification products	In the software, melt curve analysis is included in the default run method for any experiment type that uses intercalating dyes.				
of a PCR that used intercalating dyes.	1. The software plots a melt curve based on the fluorescence of the dye with respect to change in temperature.				
	<b>2.</b> Using the melt curve, the software calculates the melting temperature $(T_m)$ .				



# Workflow overview

Use the tabs across the top of the screen to navigate the workflow in the QuantStudio<sup>™</sup> desktop software.





# General procedures to set up and run experiments

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# Set global desktop software preferences

If security is enabled in the software, your account must have the permission of **Edit System Preferences**.

1. In the menu bar, select **Tools > Preferences**.

Tab	Options	Action
Defaults	Decimal Places to Show	Enter the number of significant figures in exported results.
	Language [if available]	Choose a language for the software from the dropdown menu.
Experiment	Baseline Start Cycle and Baseline End Cycle	Enter the first and last cycles to be used to calculate the baseline for runs that include amplification.
	Auto Analysis	Select to perform auto analysis at the end of each run.
	Auto Save	Select to save changes at the end of each run.
Print	Disable color when printing the Well Table	Select to disable color printing.
Data Destination	Save As Location, Export Location, and Report Location	Select where to save files and where to export results. Click <b>Browse</b> , then navigate to and select a default location.
Display Format	Date Format, Time Format, and Decimal Point Format	Select the display formats. These formats are also used in the export or import of data.

2. In the System Preferences dialog box, set preferences in each tab.

#### 3. Click Save.



# Workflow: Set up and run an experiment

Set up a template (page 16)

Create or open a template (page 17)

Enter template properties (page 17)

Confirm or edit the run method and optical filter selection (page 19)

Assign plate and well attributes (page 20)

Save a template file (page 22)

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Prepare reactions (page 23)

▼

Start and monitor a run (page 24)

For post-run procedures, see the following sections:

- Chapter 3, "General procedures to review results"
- "Export experiments or results" on page 42

# Set up a template

This section describes the general procedures to set up a template in the desktop software. For setup information for a specific experiment type, see the following sections.

- "Set up a standard curve experiment in the software" on page 47
- "Set up a relative standard curve experiment in the software" on page 57
- "Set up a comparative Ct experiment in the software" on page 58
- "Set up a genotyping experiment in the software" on page 67
- "Set up the presence/absence experiment in the software" on page 74
- "Set up a melt curve experiment in the software" on page 81
- "Set up a custom experiment in the software" on page 98

You can also use features of the desktop software to more easily set up some or all of an experiment. For example, you can set up an experiment using one of the following strategies.

- Use desktop software libraries to set up samples, targets or SNP assays, run methods, and analysis settings (see "Sample, target, and SNP assay libraries" on page 111).
- Import experiment parameters from external files or templates (see "Experiment setup using libraries, external files, and templates" on page 27).

#### Create or open a template

Create a new template or open an existing template in the **A Home** screen.

If security is enabled in the software, your account must have the permission of **Create New Template** to create a new template.

• In the 🚵 New Experiment pane, perform one of the following tasks to create a new template.

То	Action
Create a template without preexisting settings	Click Create New Experiment.
Create a template from a system template	<ul> <li>a. Select Create New Experiment &gt; Template.</li> <li>b. Navigate to and select the desired file, then click Open. System template files are installed with the software in: <drive>:\Program Files (x86)\Applied BioSystems\QuantStudio Design &amp; Analysis Software\templates, where <drive> is the drive on which the software is installed.</drive></drive></li> </ul>

- In the Some Section 2 and Secti
  - a. Click Open.
  - b. Navigate to and select the desired file, then click Open.

#### Enter template properties

If security is enabled in the software, your account must have the permission of **Edit Template Properties**.

If security is enabled in the software, your account must have the permission of **Edit Save As Destination** to select a different location to save the file.

- 1. Click the Properties tab to open and edit the experiment properties.
- 2. (Optional) In the Name field, modify the file name.

The **Name** field determines two file names:

• The initial EDT file name.

Note: *After* the initial EDT file save, modifying the **Name** field does not update the EDT file name.

To change the EDT file name *after* the initial save, click  $\Box_{\bar{*}}$  Save > Save As. In the Save As dialog box, enter the new EDT file name in the **File Name** field. Click **Browse** to edit the location to save the file, then click **Save**.

- The default file name for the EDS file created during an instrument run.
- 3. (Optional) Click the **Barcode** field, then scan or enter a plate barcode.

- 4. (Optional) Enter information in the User name field and the Comments field, if applicable.
- 5. Select an Instrument type, Block type, Experiment type, Chemistry (reagents), and Run Mode (Fast or Standard cycling) from the dropdown lists.

**Note:** The experiment type defines the available options for the template setup. For more information on the parameters defined in each experiment type, see "Experiment definitions" on page 21.

- 6. (Optional) Click Manage chemistry details (see "Enter reagent information").
- 7. (Optional) Click  $\square_{\downarrow}$  Save or click  $\square_{\downarrow}$  Save > Save as.
- 8. To save as, in the **Save As** dialog box, edit the file name in the **File Name** field, click **Browse** to edit the location to save the file, then click **Save**.

#### Enter reagent information

- 1. In the Properties tab, click Manage chemistry details.
- 2. Click + Add.
- 3. Enter the reagent type, name, part number, lot number, and expiration date.
- 4. (Optional) Add a custom attribute for a reagent.
  - a. Click + in the table header to add a column for a custom attribute.
  - b. Click the Custom Attribute column header, then enter a new attribute.
  - c. Select a cell in the Custom Attribute column, then enter its information.
  - d. (Optional) Click 🗙 in the header to delete a custom attribute from the table.
- 5. (Optional) Click  $\mathbf{X}$  to delete a reagent from the table.
- 6. Click Close.

#### Scan a barcode using the optional barcode scanner

The instrument is compatible with an optional Handheld Barcode Scanner (Cat. No. 4488442, purchased separately). The barcode scanner reads Code 128 (alphanumeric), which supports 128 ASCII character barcodes.

- 1. Click the Barcode field.
- 2. Hold the scanner 20–30 cm away from a plate or container label and aim at the center of the barcode, then press the trigger.
- 3. Slowly move the scanning beam across the barcode until the scanner emits a high-pitched tone.

When the scanner scans a barcode, it automatically transmits the following information:

- Transmits the alphanumeric equivalent of the barcode to the barcode field.
- Transmits other reagent information (Lot #, Part #, Expiration Date, etc.)

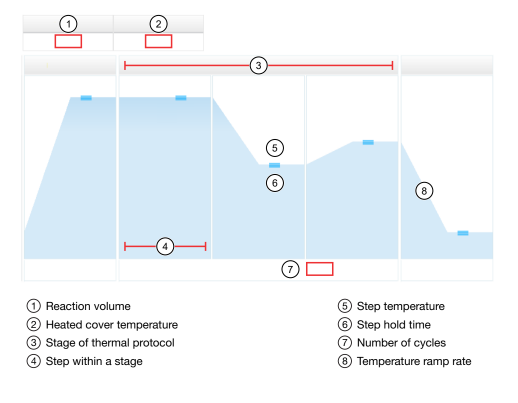
For more information about the hand-held barcode scanner, see the user documentation provided with the barcode scanner.

#### Confirm or edit the run method and optical filter selection

In the Method tab, perform the follow tasks if needed.

If security is enabled in the software, your account must have the permission of **Edit Run Method** to edit the run method.

- (Optional) Adjust the reaction volume.
- (Optional) Edit the default run method (thermal protocol).
  - The default run method is optimized for TaqMan<sup>™</sup> assays and a broad range of other reagents.
  - To edit the default run method, see "Adjust method parameters" on page 103.
- (Optional) Edit the default optical filter selection (see "Select optical filters" on page 105).
  - The default optical filter selection is for factory-calibrated (system) dyes.
  - For more information about instrument supported dyes and their calibration and optical filter selection, see the instrument user guide.



#### Method elements



#### Assign plate and well attributes using the Quick Setup subtab

If security is enabled in the software, your account must have the following permissions to perform the following tasks:

- The permission of Assign Targets / Assays to assign a target or an assay to a well
- The permission of Define New Target / Assay to add a new target or a new assay
- The permission of Assign Sample to assign a sample to a well
- The permission of Define New Sample to add a new sample
- The permission of Edit Sample to edit a sample
- The permission of **Delete Sample** to delete a sample
- The permission of Edit Passive Reference to edit the passive reference dye

Note: This section provides general procedures to set up the plate.

- For detailed procedures to set up a plate, see Appendix D, "Detailed procedures to set up plate / well details and libraries".
- For specific instructions for each experiment type, see the corresponding chapter in this guide.
- 1. In the **Plate** tab, select plate wells in the  $\parallel\parallel$  **Plate Layout** or the  $\equiv$  **Well Table**. For more information, see "Select plate wells" on page 21.
- 2. Click Quick Setup.
- 3. Assign the well attributes for the selected wells.
  - Enter the sample and target or SNP assay names in the text field.
  - Select a defined sample and target or SNP assay from the dropdown lists. For more information about defining or importing samples and targets or SNP assays, see "Define and assign well attributes (Advanced Setup subtab)" on page 107.
  - Click Advanced Setup, then change the default selections for the reporter and quencher dyes and for tasks where applicable. For more information, see "Assign a task to wells" on page 108.
- 4. (Optional) Enter comments for the selected wells.
- 5. In the Plate Attributes pane, select a Passive Reference from the dropdown list.

2

#### Select plate wells

• Select plate wells in the III Plate Layout.

То	Action
Select a single well	Click a well in the plate
Select multiple wells	Click-drag in the plate
Select contiguous wells	Shift-click wells in the plate
Select non-contiguous wells	Ctrl-click wells in the plate
Select a column of wells	Click a column header
Select all wells	Click the top-left corner of the plate grid
Select a block of wells	Click a well to define a corner, then Shift-click another well on the opposite corner
Deselect a single well	Ctrl-click the selected well

#### • Select plate wells in the $\equiv$ Well Table.

То	Action
Select a single well	Click a row in the table
Select multiple wells	Click-drag in the table
Select contiguous wells	Shift-click rows in the table
Select non-contiguous wells	Ctrl-click rows in the table
Deselect a single well	Ctrl-click the selected row

#### **Experiment definitions**

The parameters that you can define vary by experiment type.

Experiment Type	Targets	SNP Assays	Samples	Biological Replicate Groups	Passive Reference	Reference and Endogenous Controls
Standard Curve	$\checkmark$		<ul> <li>Image: A transmission of the second se</li></ul>	$\checkmark$	$\checkmark$	
Relative Standard Curve	$\checkmark$		$\checkmark$	$\checkmark$	✓	<ul> <li>✓</li> </ul>
Comparative C <sub>t</sub>	$\checkmark$		$\checkmark$	$\checkmark$	$\checkmark$	<ul> <li>✓</li> </ul>
Melt Curve	$\checkmark$		$\checkmark$		$\checkmark$	
Genotyping		$\checkmark$	$\checkmark$		$\checkmark$	
Presence / Absence	$\checkmark$		✓		$\checkmark$	
Custom	$\checkmark$		$\checkmark$	$\checkmark$	$\checkmark$	



#### Save a template file

#### Save a template file as an unlocked template

Note: You cannot save a locked template as an unlocked template.

If security is enabled in the software, your account must have the permission of **Edit Save As Destination** to select a different location to save the file.

- Save the template with the same EDT file name.
  - In any tab, click □<sub>↓</sub> Save.
  - In the menu bar, click **File Save**.
- Save the template with a new EDT file name.

Note: You cannot save a locked template with a new file name.

- In any tab, click □<sub>↓</sub> Save > Save as.
- In the menu bar, click **File Save as**.

In the **Save As** dialog box, enter the new EDT file name in the **File Name** field. Click **Browse** to edit the location to save the file, then click **Save**.

#### Save a template file as a locked template

**IMPORTANT!** Always save a backup *unlocked* version of a template before saving it as a locked template.

Record the password for the locked template because lost passwords cannot be recovered.

If security is enabled in the software, your account must have the permission of **Edit Save As Destination** to select a different location to save the file.

- 1. In the menu bar, select File > Save As Locked Template .
- 2. Enter and confirm a password, then click **OK**.

**Note:** The password is required to open the template with unlimited editing options. Without the password, a locked template can still be opened but with limited editing options.

- 3. In the Save As dialog box, edit the file name in the File Name field.
- 4. Click **Browse** to edit the location to save the file, then click **Save**.

# **Prepare reactions**

See instrument user guide for information about compatible reagents and required materials for PCR reactions.

Follow the instructions provided by the manufacturer to prepare reactions.

Follow the other guidelines described in this section.

#### Good laboratory practices for PCR and RT-PCR

- Wear clean gloves and a clean lab coat.
  - Do not wear the same gloves and lab coat that you have previously used when handling amplified products or preparing samples.
- Change gloves if you suspect that they are contaminated.
- Maintain separate areas and dedicated equipment and supplies for:
  - Sample preparation and reaction setup.
  - Amplification and analysis of products.
- Do not bring amplified products into the reaction setup area.
- Open and close all sample tubes carefully. Avoid splashing or spraying samples.
- Keep reactions and components capped as much as possible.
- Use a positive-displacement pipettor or aerosol-resistant barrier pipette tips.
- Clean lab benches and equipment periodically with 10% bleach solution or DNA decontamination solution.

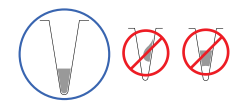
#### Guidelines for handling samples and reagents

- Use calibrated pipettors and aerosol-resistant tips.
- Prepare the reaction mixes according to the recommendations that are provided by the manufacturer of the master mixes and assay mixes.
- Include excess volume in calculations to account for the loss that occurs during reagent transfers.
- Use TE buffer or water to dilute samples and standards.
- Use care when diluting samples and standards. Mistakes or inaccuracies in making the dilutions affect data accuracy.
- Keep the dilutions and assay mix frozen and protected from light until use. Excessive exposure to light can affect the fluorescent probes or dyes.
- Perform the following tasks before each use:
  - Mix the master mix thoroughly by swirling the bottle.
  - Resuspend the assay mix by vortexing, then centrifuge the tube briefly.
  - Thaw any frozen samples by placing them on ice. When thawed, resuspend the samples by vortexing, then centrifuge the tubes briefly.



#### Guidelines for setting up the reactions in the plates or tubes

- Use good laboratory practices for PCR and RT-PCR. For more information, see "Good laboratory practices for PCR and RT-PCR" on page 23.
- Ensure that the arrangement of the PCR reactions matches the plate layout displayed in the software.
- Confirm that the liquid in each well is at the bottom of the well and free of bubbles. If it is not, centrifuge the plate again.



- Ensure that plates or tubes are properly sealed.
- Keep the reaction plate or tubes at 4°C and protected from light until you are ready to load the plate into the instrument.
- Keep the bottom of the plate clean. Fluids and other contaminants on the bottom of the plate can contaminate the sample block and cause an abnormally high background signal.
- If necessary, use a permanent marker or pen to mark a tube and the side of a plate. Do not use fluorescent markers.

# Start and monitor a run

**IMPORTANT!** Before loading a reaction-filled plate into the instrument, review the detailed procedures (see "Load and unload the plate in the instrument" on page 94).

If security is enabled in the software, your account must have the permission of Start Run.

- 1. Go to an instrument connected to the computer that is running the desktop software.
- 2. Load the plate into the instrument.



**CAUTION!** (*QuantStudio*<sup>™</sup> 3 *Real-Time PCR System and QuantStudio*<sup>™</sup> 5 *Real-Time PCR System only*) The instrument should be used by trained operators who have been warned of the moving parts hazard.

- 3. Open a template (EDT file) in the desktop software.
- 4. (*Optional*) In the **Export** tab, select **Auto Export** to export run results automatically after the run ends.
- 5. In the **Method** tab and the **Plate** tab, review the template parameters and setup.
- 6. In the Run tab, select the instrument to use from the START RUN dropdown list.

7. Accept or edit the default name for the EDS file, then click Save.

**Note:** EDS files contain the run data and results. The system creates the EDS default file name for the EDS file from the **Experiment Name** in the **Properties** tab.

- 8. During the instrument run, monitor the run:
  - In the **Run** tab of the desktop software.
  - In the instrument touchscreen.
- 9. Unload the plate from the instrument.



**CAUTION! PHYSICAL INJURY HAZARD.** During instrument operation, the plate temperature can reach 100°C. Allow it to cool to room temperature before handling.

**Note:** (*QuantStudio*<sup>™</sup> 3 *Real-Time PCR System and QuantStudio*<sup>™</sup> 5 *Real-Time PCR System only*) If the instrument does not eject the plate, contact Support.

**Note:** If the connection between the instrument and the desktop software is interrupted during the run, the instrument still completes the run. However, the run data (EDS file) must be transferred from the instrument to the desktop software using a USB drive or a network drive.

#### View real-time run information in the desktop software

You can view real-time run information in the Run tab of the desktop software.

- In the **Run** tab, view the time remaining for the run and the run status.
- In the **Run** tab, in the **Amplification Plot** subtab, view real-time data and plots.
  - Click View to select the data that are displayed in each well.
  - Select wells in the plate layout to highlight respective curves in the plot.
  - Select curves in the plot to highlight respective wells in the plate layout.

Note: For melt curve experiments, you can only monitor the melt curve plot.

#### View real-time run information on the instrument touchscreen

You can view real-time run information in the instrument home screen.

- In the instrument home screen, view the block temperature, the time remaining for the run, and the run status.
- Touch > or swipe left once to view real-time run method information.
- Touch > or swipe left twice to view real-time data and plots.



#### View real-time data and plots on the instrument touchscreen

- 1. In the instrument home screen, during an instrument run, touch > or swipe left twice.
- 2. Touch Well details.
- 3. Touch Samples, Targets, or Tasks to select a graphical representation of each selection.
- 4. Touch **Close** to return to the home screen.

#### Adjust the display of real-time plots on the instrument touchscreen

- 1. In the instrument home screen, during an instrument run, touch > or swipe left twice to view real-time data and plots.
- 2. Touch Zoom.
- **3.** Touch  $\oplus$  or  $\oplus$  to zoom in or out.
- 4. Touch the arrows to pan left, right, up, or down on the graph.
- 5. Touch Close to return to the default view.

#### View the Post-Run Summary in the desktop software

You can view a summary of the run after the run ends.

In the **Run** tab, click the **Post-Run Summary** tab to view a summary of the run, including the following information:

- Experiment Name
- User Name
- Errors Encountered
- Instrument Serial Number and Instrument Name
- Start Time, Stop Time, and Run Duration



The desktop software offers the following features so that you can more easily set up some or all of an experiment using libraries, external files, and templates.

- Use desktop software libraries to set up samples, targets or SNP assays, run methods, and analysis settings (see "Libraries overview" on page 111).
- Import some or all of an experiment setup from external files or templates (see the following table).

Option	Action	Setup information
Import sample information (define samples).	Import a sample definition file (see "Assign samples using a sample definition file" on page 99).	<ul> <li>Sample name</li> <li>(Optional) Custom sample properties</li> </ul>
Import samples, targets, and well assignments.	Import a plate setup file (see "Assign samples and targets using plate setup files" on page 100).	<ul> <li>Plate setup information:</li> <li>Well number</li> <li>Sample name</li> <li>Sample color</li> <li>Target name</li> <li>Dyes</li> <li>(Optional) Other well information</li> </ul>
Set up the plate layout in a spreadsheet without saving to a special format. <i>or</i> Use a subset of the columns in a plate layout spreadsheet.	Copy-paste from an XLS file (see "Assign targets, samples, and biological replicate groups from an XLS file" on page 101).	<ul> <li>Plate setup information:</li> <li>Well number</li> <li>Sample name</li> <li>Biological Group</li> <li>Target name</li> <li>Task</li> <li>Dyes</li> <li>Quantity</li> <li>Comments</li> <li>(Optional) Other well information</li> </ul>
Use a complete template setup from an existing EDT or EDS file.	Create a new template from an existing template or run results file (see "Create new EDT files using existing EDT and EDS files" on page 102).	<ul> <li>Plate setup information, as above</li> <li>Reagent information</li> <li>Thermal protocol</li> <li>Analysis settings</li> </ul>

#### Table 1 Experiment setup from external files or templates



# General procedures to review results

About the quantification cycle (Cq)	28
Overview of the Results tab	29
Workflow: General procedures to review the run results	31
Guidelines for viewing and analyzing results	31
Assess results in the Amplification Plot	32
Assess results in the Well Table view	37
Review the dye signal profile using the Multicomponent Plot	38
Review the signal profile using the Raw Data Plot	39
Review the flags in the QC Summary	40
View calibration results in the desktop software	41
Export experiments or results	42

This section includes information about reviewing results and configuring analysis settings for all experiment types. For information about a specific experiment type, see the corresponding chapter in this guide.

For step-by-step instructions for results review procedures, see the desktop software Help.

# About the quantification cycle (Cq)

Term	Name	Description
Cq	Quantification cycle	$C_q$ is the general form for gene expression metrics. $C_q$ values (both $C_t$ and $C_{rt}$ ) are used as the primary input values for sample quantification experiments: absolute quantification (AQ) and relative quantification (RQ).
		$C_t$ and $C_{rt}$ are the algorithm-specific calculations of $C_q$ .
Ct	Threshold cycle	The PCR cycle number at which the fluorescence signal meets the threshold in the amplification plot.
		$C_{t}$ is the gene expression metric result when using the Baseline Threshold Algorithm.
C <sub>rt</sub>	Relative threshold cycle	The PCR cycle number for the threshold calculated from the modeled amplification efficiency profile.
		$C_{rt}$ is the gene expression metric result when using the Relative Threshold Algorithm.

# 3

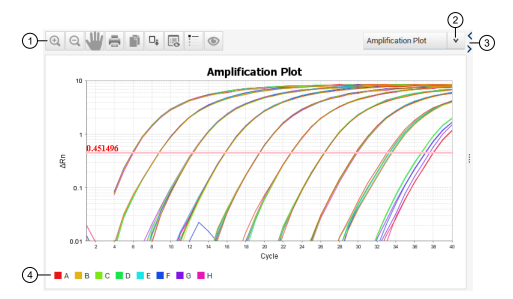
# Overview of the Results tab

Review and analyze run data in the **Results** tab. In the **Results** tab, two additional tools display at the right of the workflow bar.

Properties Method Plate Run	Results Export	Analyze 🔅	
-----------------------------	----------------	-----------	--

- Click Analyze after omitting wells or changing the analysis settings.
- Click to access analysis settings.

Note: The analysis settings and plots that are available vary by experiment type.



#### Figure 1 Plot pane

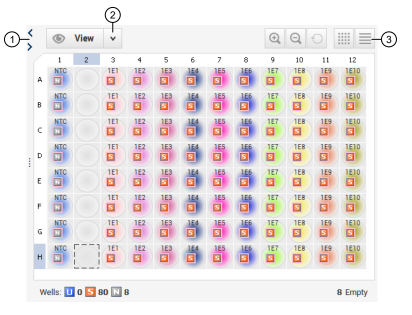
- 1 Plot toolbar
- 2 Plot selection list (varies by experiment)

The Plot toolbar includes the following options:

- Zoom in and out
- Print or copy plot image
- Save plot as image file

- ③ Expand/contract the plot pane display
- ④ Plot legend
- Configure plot properties
- Show/hide plot legend
- Configure plot settings

3



#### Figure 2 Plate Layout

- (1) Expand/contract the Plate Layout display (layout is expanded in this figure)
- 2 View: Select well properties to display
- 3 Plate Layout toolbar

The Plate Layout toolbar includes the following options:

- · Zoom in and out
- · Fit plate to window

- Display Plate Layout
- Display Well Table

			2		3				
1-	Solution State		Group by	*		$+ \bigcirc \qquad \blacksquare = -4$			
	#	Well	Sample Na	Flag	Target Na	Task	Dyes	Ст	
	1	A1	NTC	•	KAZ	NTC	FAM-NF	Undeter	0
	2	A2		•					
	3	A3	1E1	1	KAZ	STAND	FAM-NF	38.028	
:			150		1/17	OTAND		00.056	

#### Figure 3 Well Table

- (1) Expand/contract the **Well Table** display (table is expanded in this figure)
- 2 View: Select well properties to display
- ③ Group by: Select a parameter by which to group well rows
- (4) Well Table toolbar

The Well Table toolbar includes the following options:

- Expand grouped rows
- Collapse grouped rows

- Display Plate Layout
- Display Well Table

# Workflow: General procedures to review the run results

When a run is complete, the desktop software automatically analyzes the run data using the analysis settings that are specified during template development. The software then displays the run results in the **Results** tab.

View the Amplification Plot to confirm or correct threshold and baseline settings (page 32)

#### ▼

Assess the experiment plot for the experiment (for example, view the Allelic Discrimination Plot for genotyping experiments)

(see the corresponding chapter in this guide)

▼

Review data for outliers and *(optional)* omit wells (page 35)

▼

(Optional) View the Multicomponent Plot to review the dye signal profile (page 38)

▼

(Optional) View the Raw Data Plot to review the signal profile (page 39)

(Optional) Review flags in the QC Summary (page 40)

▼

(Optional) Configure the analysis settings (page 115)

**IMPORTANT!** If you omit wells or configure the analysis settings, click **Analyze** to reanalyze the data.

### Guidelines for viewing and analyzing results

- For information about adjusting the views in the **Results** tab, see the desktop software Help.
- To reanalyze the data, select all the wells in the III Plate Layout, then click Analyze.
- To enable auto-analysis of data after a run, select Tools > Preferences > Experiment, then select Auto Analysis.

# Assess results in the Amplification Plot

#### **Amplification Plot overview**

The Amplification Plot displays sample amplification as a function of cycle number or well. You can use the amplification plot to perform the following tasks:

- Confirm or correct baseline and threshold values.
- Identify outliers.
- Identify and examine abnormal amplification. Abnormal amplification can exhibit one of the following characteristics:
  - Increased fluorescence in negative control wells.
  - Absence of detectable fluorescence at an expected cycle.

**Note:** If you notice abnormal amplification or a complete absence of fluorescence, see the instrument user guide for troubleshooting information.

Three plots are available. Some plots can be viewed as a linear or log<sub>10</sub> graph.

#### Table 2Amplification Plot types

Plot type	Description	Use to
∆Rn vs Cycle	$\Delta Rn$ is the magnitude of normalized fluorescence signal, relative to the baseline fluorescence, generated by the reporter at each cycle during the PCR amplification.	<ul> <li>Identify and examine irregular amplification.</li> <li>View threshold values for the run.</li> </ul>
Rn vs Cycle	Rn is the fluorescence signal from the reporter dye normalized to the fluorescence signal from the passive reference.	<ul> <li>Identify and examine irregular amplification.</li> <li>View baseline values for the run.</li> </ul>
C <sub>t</sub> vs Well	C <sub>t</sub> is the PCR cycle number at which the fluorescence meets the threshold in the amplification plot.	<ul> <li>Locate outlying amplification (outliers).</li> </ul>

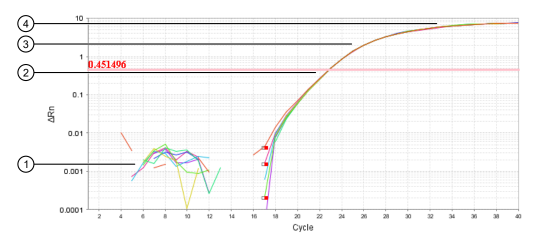
#### Assess the overall shape of the Amplification Plot curves

You can assess the overall shape of the Amplification Plot curves in the **Results** tab.

If no data are displayed in the **Results** tab, click **Analyze**.

- 1. In the **Results** tab, select **Amplification Plot** from the dropdown list.
- 2. Click ( to configure the plot, then make the following selections:
  - Plot Type: ΔRn vs Cycle
  - Graph Type: Log
  - Plot Color: Target, Sample, or Well

The Amplification Plot is displayed for the selected wells in the III Plate Layout.





A typical amplification curve has four distinct sections:

① Baseline③ Linear phase② Exponential (geometric) phase④ Plateau phase

#### Confirm or correct threshold settings

- 1. In the Results tab, select Amplification Plot from the dropdown list.
- 2. Click ( to configure the plot, then make the following selections:
  - Plot Type: ΔRn vs Cycle
  - Graph Type: Log
  - Plot Color: Target, Sample, or Well

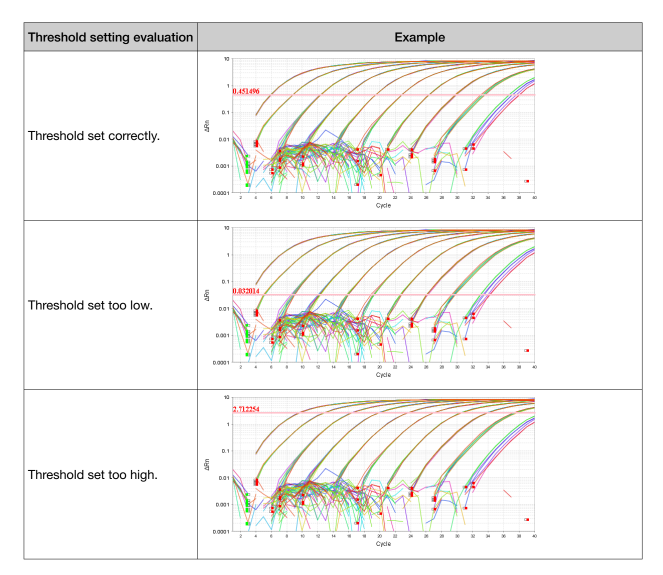
The Amplification Plot is displayed for the selected wells in the III Plate Layout.

- 3. (Optional) Adjust the threshold.
  - Click-drag the threshold bar into the exponential phase of the curve.
  - Configure the C<sub>t</sub> analysis settings (see "Ct settings overview" on page 116).

#### Table 3 Examples of threshold settings

Set the threshold in the exponential phase of the amplification curve. A threshold set above or below the optimum can increase the standard deviation of the replicate groups.

QuantStudio™ Design and Analysis Desktop Software User Guide



#### Confirm or correct baseline settings

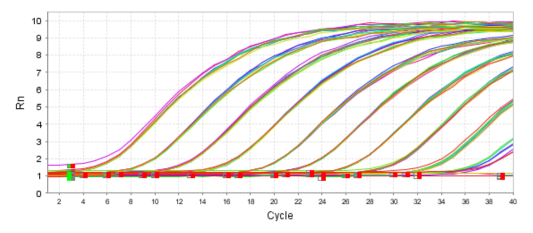
- 1. In the Results tab, select Amplification Plot from the dropdown list.
- 2. Click  $\circledast$  to configure the plot, then make the following selections:
  - Plot Type: Rn vs Cycle
  - Graph Type: Linear
  - Plot Color: Well
  - Select Show: Baseline Start / Baseline End

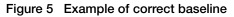
Note: The start and end cycles are used to calculate the baseline.

The Amplification Plot is displayed for the selected wells in the III Plate Layout.

The start  $(\blacksquare)$  and end  $(\blacksquare)$  cycles display for each well.

**3.** (*Optional*) Adjust the start and end cycle values for the baseline (see "Ct settings overview" on page 116).





Set the end cycle ( ) a few cycles before the cycle number where significant fluorescence signal is detected.

#### Omit outliers from analysis

Outlier wells have  $C_q$  ( $C_t$  or  $C_{rt}$ ) values that differ significantly from the average for the associated replicate wells. To ensure  $C_q$  ( $C_t$  or  $C_{rt}$ ) precision, consider omitting the outliers from analysis.

If security is enabled in the software, your account must have the permission of **Omit Wells for Analysis**.

- 1. In the Results tab, select Amplification Plot from the dropdown list.
- 2. Click (1), then make the following selections to configure the plot:
  - Plot Type: Ct vs Well
  - Graph Type: Linear
  - Plot Color: Well

The C<sub>t</sub> values are displayed for the selected wells in the  $\blacksquare$  Plate Layout.

- 3. Click  $\equiv$  to examine the Well Table for outliers.
  - a. Select Group by > Replicate.
  - b. Identify outliers in each replicate group.
     Outlier wells typically have one or more QC flags.



- 4. Omit outliers in either the  $\equiv$  Well Table or  $\equiv$  Plate Layout view.
  - In the  $\equiv$  Well Table, select Omit in outlier rows of the table.
  - In the I Plate Layout, right-click a well, then select Omit.
- 5. Click Analyze to reanalyze the run data with any outliers removed.

#### Optimize display of negative controls in the Amplification Plot

- 1. In the **Results** tab, select **Amplification Plot** from the dropdown list.
- 2. Click 
   to configure the plot, then make the following selections:
  - Plot Type: ΔRn vs Cycle
  - Graph Type: Linear
  - Plot Color: Target
  - Deselect Show: Threshold
  - Deselect Show: Baseline Start / Baseline End
- 3. In either the  $\parallel\parallel$  Plate Layout or  $\equiv$  Well Table, select the negative control wells (wells that should not have amplification for a particular target).
- 4. Click 🔜 (configure plot properties), then select the **Y** Axis tab.
  - a. Deselect Auto-adjust range.
  - b. Enter Minimum value of -1.
  - c. Enter Maximum value of 2.
  - d. Click Save.

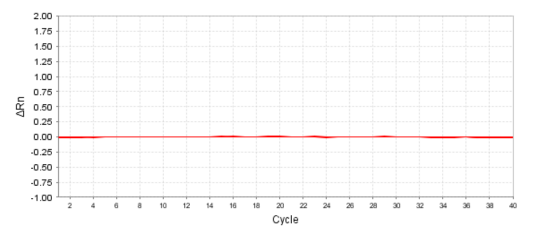


Figure 6 Example Amplification Plot of negative controls

The linear plot displays the Amplification Plot for negative controls as smooth lines. The expanded Y axis displays low levels of amplification.



## Assess results in the Well Table view

#### Well Table overview

The  $\equiv$  **Well Table** displays data for each well in the reaction plate. The data that are displayed depend on the specific experiment type and can include the following information:

- Sample name, target name, task, and dyes
- Values that are specific to particular stage of the method
   For example: C<sub>t</sub> or C<sub>rt</sub>, normalized fluorescence (Rn), or melt temperature (T<sub>m</sub>)
- Values that are specific to a particular experiment type For example: genotype calls, presence/absence calls, or quantities
- Omitted wells
- QC flags
- Comments

## Group or sort the Well Table

Some of the possible options for grouping or sorting the  $\equiv$  **Well Table** are described in the following table. Available grouping categories depend on the specific experiment type and analysis settings.

If the data are grouped, this order is retained in the report (see "Print a report of run data and results" on page 44).

**Note:** You can select multiple columns when sorting, but you can only make one selection for grouping rows.

Group category	Description	Notes
Replicate <sup>[1,2,3]</sup>	Grouped by replicate	<ul> <li>Examine the C<sub>t</sub> or quantity values for each replicate group to assess the precision of C<sub>t</sub> values.</li> </ul>
Flag	Grouped as flagged and unflagged wells	<ul> <li>A flag indicates that the software found a potential error in the flagged well.</li> <li>For more information about QC flags, see the desktop software Help.</li> </ul>
Ct <sup>[1,2,4]</sup>	Grouped by C <sub>t</sub> value	<ul> <li>C<sub>t</sub> value &lt; 8—There may be too much template in the reaction.</li> <li>C<sub>t</sub> value &gt; 35—There may be a low amount of target in the reaction; for C<sub>t</sub> values &gt; 35, expect a higher standard deviation.</li> </ul>
RQ <sup>[2]</sup>	Grouped by RQ value	<ul> <li>RQ value &lt; 1—There is less relative target in the test sample as compared to the calibrator sample.</li> <li>RQ value &gt; 1—There is more relative target in the test sample as compared to the calibrator sample.</li> </ul>
<b>Call</b> <sup>[4,5]</sup>	Grouped by genotype call or presence/absence call	_



#### (continued)

Group category	Description	Notes
Tm1 <sup>[3]</sup>	Grouped by T <sub>m</sub> value	Tm1 refers to the dominant peak.
		This grouping category is only applicable for the following experiment types:
		Melt curve experiments
		<ul> <li>Any experiment with a melt curve data collection step (e.g., absolute standard curve)</li> </ul>

<sup>[1]</sup> For standard curve experiments.

<sup>[2]</sup> For relative standard curve and comparative  $C_t$  ( $\Delta\Delta C_t$ ) experiments.

<sup>[3]</sup> For melt curve experiments.

<sup>[4]</sup> For genotyping experiments.

<sup>[5]</sup> For presence/absence experiments.

# Review the dye signal profile using the Multicomponent Plot

## **Multicomponent Plot overview**

The Multicomponent Plot displays the complete spectral contribution of each dye over the duration of the PCR run.

Use the Multicomponent Plot to obtain the following information.

- Confirm that the signal from the passive reference dye remains unchanged throughout the run.
- Review reporter dye signal for spikes, dips, and/or sudden changes.
- Confirm that no amplification occurs in the negative control wells.

## View and assess the Multicomponent Plot

You can view and assess the Multicomponent Plot in the Results tab.

If no data are displayed in the Results tab, click Analyze.

- 1. In the Results tab, select Multicomponent Plot from the dropdown list.
- 2. Click 
   to configure the plot, then make the following selections:
  - Plot Color: Dye

The Multicomponent Plot is displayed for the selected wells in the III Plate Layout.

3

3. In the IIII Plate Layout, select wells one at a time, then examine the Multicomponent Plot for the following plot characteristics.

Plot characteristic Description		
Passive reference dye	The passive reference dye fluorescence signal should remain relatively constant throughout the PCR process.	
Reporter dye	The reporter dye fluorescence signal should display a flat region corresponding to the baseline, followed by a rapid rise in fluorescence as the amplification proceeds.	
Irregularities in the signal Spikes, dips, and/or sudden changes in the fluorescence signal may have an impatheet the data.		
Negative control wells	The negative control wells should show no significant increase in fluorescence signal.	

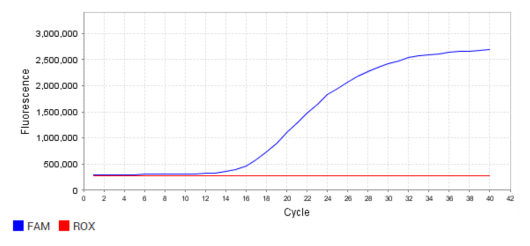


Figure 7 Example Multicomponent Plot (single well)

## Review the signal profile using the Raw Data Plot

## Raw Data Plot overview

The Raw Data Plot displays the raw fluorescence signal (not normalized) for each optical filter during each cycle of the real-time PCR.

View the Raw Data Plot to confirm a stable increase in signal (no abrupt changes or dips) from the appropriate filter.



## View and assess the Raw Data Plot

You can view and assess the Raw Data Plot in the Results tab.

If no data are displayed in the Results tab, click Analyze.

- In the Results, select Raw Data Plot from the dropdown list. The Raw Data Plot is displayed for the selected wells in the IIII Plate Layout.
- 2. Click ( to display the Show Cycle scale.
- **3.** Click-drag the **Show Cycle** pointer from cycle 1 to cycle 40, and confirm that each filter displays the characteristic signal increase.

For more information on each filter set, see the instrument user guide (see Appendix F, "Documentation and support").



Figure 8 Example Raw Data Plot

## **Review the flags in the QC Summary**

The **QC Summary** in the **Results** tab displays a list of the QC flags, including the flag frequency and location.

If no data are displayed in the **Results** tab, click **Analyze**.

- 1. In the Results tab, select QC Summary from the dropdown list.
- 2. Review the Flag Details table and the summary.

The Wells column of the Flag Details table identifies wells that triggered a flag.

**3.** (*Optional*) In the **Flag Details** table, click each flag to display a brief description of the flag. For more information about the flag, see the desktop software Help.

## View calibration results in the desktop software

Calibration results obtained on the instrument can be transferred via USB to the desktop software.

- 1. In the 🕂 Home screen, click 🔬 Open.
- 2. Navigate to and select the desired calibration EDS file.

If you are viewing calibration data files,  $\mathcal{M}$  Calibration QC Status is displayed to the right of the **Hate Layout** and **E Well Table**.

Note: For more information about using the  $\parallel\parallel$  Plate Layout and  $\equiv$  Well Table, see Chapter 3, "General procedures to review results".

#### **Review ROI/Uniformity calibration results**

- 1. In the ROI tab, select a Filter Set from the dropdown list to see the corresponding results.
- 2. In the Uniformity tab, use the IIII Plate Layout, ≡ Well Table, and M Calibration QC Status to review the run results.

#### **Review Background calibration results**

- 1. Select the plate wells in the **IIII** Plate Layout or the **E** Well Table to view the corresponding curves. For more information, see "Select plate wells" on page 21.
- **2.** Review data in the  $\equiv$  **Well Table**.
  - a. Review the results for each well in tabular format.
  - b. Sort the wells according to well or normalized fluorescence with each filter.
  - c. Select wells to review data in the analysis plot.
- 3. Click *A* Calibration QC Status to review the quality of the calibration data.

#### **Review Dye calibration results**

- 1. Select a Dye row in the **Calibration** table to view the corresponding analysis data plot.
- 2. Select the plate wells in the **IIII** Plate Layout or the **IIII** Well Table to view the corresponding curves. For more information, see "Select plate wells" on page 21.
- **3.** Review data in the  $\equiv$  **Well Table**.
  - a. Review the results for each well in tabular format.
  - b. Sort the wells according to well or normalized fluorescence with each filter.
  - c. Select wells to review data in the analysis plot.
- 4. Click A Calibration QC Status to review the quality of the calibration data.



## Override the calibration data

Each EDS file contains the calibration data from the instrument on which it was run. You can use calibration data from another instrument for analysis of your run data.

- Calibration data must be from the same block type (96-well 0.2-mL block, 96-well 0.1-mL block, or 384-well block).
- Calibration data for a QuantStudio<sup>™</sup> 1 Real-Time PCR Instrument run must be from a QuantStudio<sup>™</sup> 1 Real-Time PCR Instrument run.
- Calibration data from a QuantStudio<sup>™</sup> 5 Real-Time PCR Instrument run can override calibration data for a QuantStudio<sup>™</sup> 3 Real-Time PCR Instrument run.
- Calibration data from a QuantStudio<sup>™</sup> 3 Real-Time PCR Instrument run can *not* override calibration data for a QuantStudio<sup>™</sup> 5 Real-Time PCR Instrument run.
- 1. Open the EDS file to recalibrate.
- 2. Select Analysis > Override Calibration > Use Calibration From Another File.
- 3. Navigate to, then select the EDS file containing the alternative calibration data.
- 4. Click Open.
- 5. (*Optional*) To revert to the original calibration data, select **Analysis ► Override Calibration ► Revert To Original Calibration**.

## Export experiments or results

For step-by-step instructions for exporting experiments or results, see the desktop software Help.

## Options for exporting the plots

То	Action	
Save a plot as an image file	In the Results tab, click . (Save to image file).	
	The 📭 (Save to image file) button is located above the plot.	
	Use the dropdown list above the plot to select a plot type.	
Print a plot	Click 📇 (Print).	
	The 🚍 (Print) button is located above the plot.	
	Use the dropdown list above the plot to select a plot type.	
Copy a plot to the clipboard	Click 👔 (Copy).	
	The 👔 (Copy) button is located above the plot.	
	Use the dropdown list above the plot to select a plot type.	

#### Export run data results

You can customize the exported results. See "Customize exported results" on page 43.

If security is enabled in the software, your account must have the permission of **Edit Export Destination** to edit the location in the **Export Location** field.

- 1. Select the Export tab.
- (Optional) Edit the file name in the File Name field.
   The default file name is set up in the Name filed of the Properties tab.
- 3. Select a file type from the File Type dropdown list.
  - QuantStudio
  - RDML

For more information, see "Export configurations" on page 45.

- **4.** If **QuantStudio** was selected as the file type, select a file format from the dropdown list. For more information, see "Export configurations" on page 45.
- 5. Select a location to save the file in the **Export Location** field.
- 6. Select the **Open exported files when complete** checkbox if you would like to open the file immediately.
- 7. In the **Content** pane, select the checkboxes that are associated with the information to include in the export.

Selecting the content is not available if **RDML** was selected as the file type.

- 8. In the **Options** pane, select the radio button to include all of the data in a single file or to export separate files.
  - Unify the above content into one file
  - Split the above content items into individual files

For a XLS file, the content is separated onto different tabs if one file is selected. For a TXT file, the content is included sequentially in the file if one file is selected. The file names are appended with the content type if individual files are selected.

9. Click Export.

#### Customize exported results

Ensure that the content is selected. The options that are selected determine what is customized.

1. In the Export tab, click Customize.

The **Customize** dialog box is displayed. Each tab in the dialog box corresponds to content item that was selected.

2. In the **Customize** dialog box, select the tab that corresponds to the content to customize.



- 3. In the Select Content pane, select items to include in the exported file.
- 4. In the **Results** tab, select the following checkboxes to skip the corresponding data from being exported.
  - Skip Empty Wells checkbox
  - Skip Omitted Wells checkbox
- 5. Click Close to return to the Export tab.

#### Print the plate layout

- 1. From any tab, click **File > Print**.
- 2. In the Print Plate Layout dialog box, select the radio button that corresponds with the color.
  - Use sample color radio button
  - Use task color radio button
- 3. Click **Print** to open the system dialog box for the printer.
- 4. Click **Preview** to open the **Print Preview** dialog box. A preview of the plate layout is displayed.
- 5. In the **Print Preview** dialog box, click one of the following options.
  - Click Save As PDF to save the file.
  - Click **Print Report** to open the system dialog box for the printer.

## Send run results to PowerPoint™

- 1. Open an EDS file.
- 2. In the menu bar, click File > Send to PowerPoint.
- 3. In the Create Slides dialog box, select the content to include in the slides, then click Apply.

**Note:** Analysis plots are not available for slides until the run status is complete and the run data are analyzed.

## Print a report of run data and results

If security is enabled in the software, your account must have the permission of **Edit Report Destination** to select a different location to save the file.

- 1. Open an EDS file.
- 2. In the menu bar, select File > Print Report.
- Select the run data and results to include in the report.
   The available data and results vary according to the experiment type.

If the data were grouped by a specific parameter, that order is retained in the report. For information about grouping, see Figure 3 on page 30 and "Group or sort the Well Table" on page 37.

- 4. Click **Print Report** to open the system dialog box for the printer.
- 5. (Optional) Click Print Preview.

A preview of the report is displayed in the **Print Preview** dialog box.

- 6. In the **Print Preview** dialog box, click one of the following options.
  - Click Save As PDF to save the file.
  - Click **Print Report** to open the system dialog box for the printer.
- 7. In the Save As dialog box, edit the file name in the File Name field.
- 8. Click **Browse** to edit the location to save the file, then click **Save**.

## **Export configurations**

Data type	Description	File format
Plate setup files, for future experiments	Plate setup information For example, the well number, sample name and color, target name, dyes, and other reaction plate contents.	<ul><li>XLS</li><li>XLSX</li><li>TXT</li></ul>
Analyzed data, for further analysis	QuantStudio™ format	<ul><li>XLS</li><li>XLSX</li><li>TXT</li></ul>
	RDML (Real-Time PCR Data Markup Language) format Used for standard curve, relative standard curve, and comparative $C_t$ experiments.	RDML



# Set up, run, and review standard curve experiments

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## Standard curve experiments

## Overview

Use standard curve experiments to determine absolute target quantity in samples.

In a standard curve experiment, the software performs the following tasks.

- 1. The software measures amplification of the target in a standard dilution series and in test samples.
- 2. The software generates a standard curve using data from the standard dilution series.
- 3. The software uses the standard curve to interpolate the absolute quantity of target in the test samples.

## **Reaction types**

Multiple targets can be assayed in a standard curve experiment, but each target requires its own standard curve.

Table 4 Reaction types for standard curve experiments

Reaction type (task)	Sample description	
Standard	A sample that contains known or known relative quantities of the target	
	<ul> <li>For known quantities—Quantify the target in the standard sample using an independent method.</li> </ul>	
	• For known relative quantities—Generate a relative dilution series of the target standards.	

Reaction type (task)     Sample description	
Unknown	Test sample
No-template control	Water or buffer
(NTC/ Negative Control)	No amplification of the target should occur in NTC wells.

#### Table 4 Reaction types for standard curve experiments (continued)

- The precision of quantification experiments improves as the number of replicate reactions increases. Set up the number of replicates appropriate for the experiment.
- For accurate and precise efficiency measurements, set up the standard dilution series with at least five dilution points over a broad range of standard quantities, 4 to 6 logs (10<sup>4</sup>- to 10<sup>6</sup>-fold). A concentrated template, such as a plasmid or PCR product, is best for this purpose.
   A narrow range of standard quantities may be appropriate if the amount of standard is limited, the target is in low abundance, or the target is known to fall within a given range.

#### **Compatible PCR options**

 Table 5
 PCR options for standard curve experiments

Single- or multiplex PCR	PCR or RT-PCR <sup>[1]</sup>	Detection chemistry
Singleplex	PCR	TaqMan™
Multiplex	1-step RT-PCR	SYBR™ Green
	2-step RT-PCR	

<sup>[1]</sup> RT-PCR: reverse transcription-PCR

## Set up a standard curve experiment in the software

- 1. In the **Home** screen, create or open a template.
  - In the 🚵 New Experiment pane, click Create New Experiment to create a new template.
  - In the Some Dependence of the pane, click Open to select and open an existing template.
- 2. In the **Properties** tab, enter the template information.
- In the Method tab, adjust the reaction volume.
   For most experiments, the default run method is appropriate.
- 4. In the **Plate** tab (**Quick Setup**), assign plate attributes.
  - a. In the Plate Attributes pane, select the Passive Reference from the dropdown list.
- 5. In the Plate tab (Quick Setup), define and assign well attributes.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.



- b. Assign samples and targets to selected wells.
  - Enter new sample and target names in the text fields.
  - Select previously defined samples and targets from the dropdown lists.

**Note:** New sample or target names entered in the **Quick Setup** subtab are automatically populated with default values for **Reporter** (FAM) and **Quencher** (NFQ-MGB) and assigned a task (U **Unknown**). Edit these values in the **Advanced Setup** subtab.

- 6. (*Optional*) In the **Plate** tab, set up standard dilutions (see "Define and set up standard dilutions" on page 109).
- 7. (Optional) In the Plate tab (Advanced Setup), assign tasks.
  - a. Select wells in the iiii Plate Layout or the  $\equiv$  Well Table.
  - **b.** In the **Targets** table, select the checkbox of a target, then select a task from the dropdown list.

Reaction type	Task
Unknown (test sample)	U
No-template control	N

8. (*Optional*) In the **Plate** tab (**Advanced Setup**), define and assign biological replicate groups (see "Define and assign biological replicate groups" on page 110).

In the **Plate** tab (**Advanced Setup**), ensure the **Samples** table contains the following sample information:

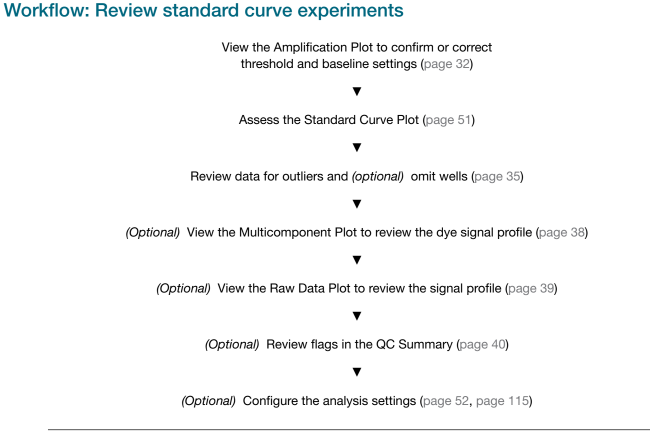
- One sample name for each technical replicate group of an unknown sample
- (Optional) A standard sample for each target

## Set up and run the PCR reactions

- 1. Assemble the PCR reactions. Follow the manufacturer's instructions for the reagents and follow the plate layout set up in the software (see "Prepare reactions" on page 23).
- 2. In the desktop software, open the appropriate template (EDT file).
- **3.** Load the reaction plate into the instrument and start the run (see "Start and monitor a run" on page 24).

## 4

## **Review results**



**IMPORTANT!** If you omit wells or configure the analysis settings, click **Analyze** to reanalyze the data.



## Standard Curve Plot overview

The Standard Curve Plot displays the standard curve for samples designated as standards. The software calculates the quantity of an unknown target from the standard curve.

Table 6 Results or metrics to review in the Standard Curve Plo	Table 6	Results or metrics	s to review in the	Standard Curve Plot
--	---------	--------------------	--------------------	---------------------

Results or metrics	Description	Criteria for evaluation
Slope and amplification efficiency	The amplification efficiency is calculated using the slope of the regression line in the standard curve.	<ul> <li>A slope close to -3.3 indicates optimal, 100% PCR amplification efficiency.</li> <li>Factors that affect amplification efficiency: <ul> <li>Improper design of the primer and probe</li> </ul> </li> <li>Range of standard quantities—For accurate and precise efficiency measurements, use a broad range of standard quantities, 5 to 6 logs (10<sup>5</sup>- to 10<sup>6</sup>-fold).</li> <li>Number of standard replicates—For accurate efficiency measurements, include replicates to decrease the effects of pipetting inaccuracies.</li> <li>PCR inhibitors—PCR inhibitors and contamination in the reaction can reduce amplification efficiency.</li> <li>Other possible factors: <ul> <li>Component and properties of the reaction mix, such as salt content, DMSO, pH, etc.</li> <li>Inaccurate sample or reagent pipetting</li> <li>Improper analysis settings</li> <li>Incorrect plate setup</li> </ul> </li> </ul>
R <sup>2</sup> value (correlation coefficient)	The $R^2$ value is a measure of the closeness of fit between the regression line and the individual $C_q$ data points of the standard reactions.	<ul> <li>A value of 1.00 indicates a perfect fit between the regression line and the data points.</li> <li>An R<sup>2</sup> value &gt; 0.99 is desirable.</li> </ul>
Error	The standard error of the slope of the regression line in the standard curve. The error can be used to calculate a confidence interval (CI) for the slope and therefore the amplification efficiency.	Acceptable value is determined by the experimental criteria.

Results or metrics	Description	Criteria for evaluation
C <sub>t</sub> values	The threshold cycle $(C_t)$ is the PCR cycle number at which the fluorescence level meets the threshold.	<ul> <li>A C<sub>t</sub> value &gt; 8 and &lt; 35 is desirable.</li> <li>C<sub>t</sub> value &lt; 8—There may be too much template in the reaction.</li> <li>C<sub>t</sub> value &gt; 35—There may be a low amount of target in the reaction; for C<sub>t</sub> values &gt; 35, expect a higher standard deviation.</li> </ul>

Table 6 Results or metrics to review in the Standard Curve Plot (continued)

## View and assess the Standard Curve Plot

You can view and assess the Standard Curve Plot in the Results tab.

If no data are displayed in the **Results** tab, click **Analyze**.

- 1. Select Standard Curve from the dropdown list.
- 2. Click 
   to configure the plot, then make the following selections:
  - Target: Select the target of interest
  - Plot Color: Sample, Target, or Task
  - Select all wells in the III Plate Layout

The Standard Curve Plot is displayed. The slope, R<sup>2</sup> value, amplification efficiency, and error are displayed below the plot.

- 3. Confirm that the slope, R<sup>2</sup> value, amplification efficiency, and error meet the experimental criteria.
- 4. Visually check that all unknown sample C<sub>q</sub> (C<sub>t</sub> or C<sub>rt</sub>) values fall within the standard curve range.
- 5. In the  $\equiv$  Well Table, use the Group By dropdown list to confirm that the C<sub>q</sub> (C<sub>t</sub> or C<sub>rt</sub>) values of all replicate samples meet the experimental criteria.

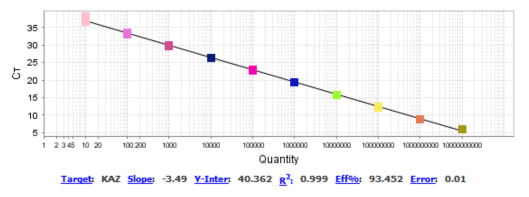


Figure 9 Example Standard Curve Plot



If the results do not meet the experimental criteria, troubleshoot using one of the following strategies:

- Omit wells, then reanalyze. For more information, see "Omit outliers from analysis" on page 35.
- Repeat the experiment, adjusting the template setup and analysis settings to improve results.

To learn more about the Standard Curve Plot, see "Standard Curve Plot overview" on page 50.

#### Standard curve settings overview

You can use the standard curve from another experiment and apply it to the current experiment. The two experiments must be from the same instrument type, block type, and run method.

To import an external standard curve, select **Analysis Settings** > Standard Curve Settings, then follow the instructions on the screen.

For step-by-step instructions for adjusting the standard curve settings, see the desktop software Help.



# Set up, run, and review relative standard curve experiments and comparative $C_t$ experiments

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## **Relative standard curve experiments**

## Overview

Use relative standard curve experiments to determine relative target quantity in samples.

In a relative standard curve experiment, the software performs the following tasks.

1. The software measures amplification of the target of interest and of an endogenous control target in a standard dilution series, in a reference (calibrator) sample, and in test samples.

The endogenous control is a target that is expressed equally in all samples; examples of endogenous controls are  $\beta$ -actin, GAPDH, and 18S ribosomal RNA. The software can algorithmically incorporate multiple endogenous control targets in relative quantification calculations.

The reference sample is used as the basis for relative quantification results (or  $1 \times$  sample). For example, in a study of drug effects on gene expression, an untreated control is an appropriate reference sample.

- 2. The software generates standard curves for the target of interest and the endogenous control using data from the corresponding standard dilution series.
- 3. The software uses the standard curves to interpolate the quantities of the target of interest and the endogenous control in each sample. The target quantity in each sample is then normalized to the sample's endogenous control quantity.
- 4. To determine the relative quantity of the target in test samples, the software divides the normalized target quantity in the sample by the normalized target quantity in the reference sample.

For a comparison of this method to the comparative  $C_t (\Delta \Delta C_t)$  method, see "Relative quantitation: relative standard curve vs. comparative Ct" on page 56.

## **Reaction types**

Relative standard curve experiments include the following reaction types for the endogenous control target and each target of interest.

Reaction type (task)	Sample description	
Standard	A sample that contains known or known relative quantities of the target	
	<ul> <li>For known quantities—Quantify the target in the standard sample using an independent method.</li> </ul>	
	<ul> <li>For known relative quantities—Generate a relative dilution series of the target standards.</li> </ul>	
Reference sample <sup>[1]</sup>	The sample that is used as the basis for relative quantification results	
Unknown	Test or reference sample	
No-template control (NTC/ Negative Control)	Water or buffer	
	No amplification of the target should occur in NTC wells.	

 Table 7 Reaction types for relative standard curve experiments

<sup>[1]</sup> To identify a sample as a reference sample, review the relative quantification settings.

• The precision of quantification experiments improves as the number of replicate reactions increases. Set up the number of replicates appropriate for the experiment.

• For accurate and precise efficiency measurements, set up the standard dilution series with at least five dilution points over a broad range of standard quantities, 4 to 6 logs (10<sup>4</sup>- to 10<sup>6</sup>-fold). A concentrated template, such as a plasmid or PCR product, is best for this purpose.

A narrow range of standard quantities may be appropriate if the amount of standard is limited, the target is in low abundance, or the target is known to fall within a given range.

## **Compatible PCR options**

#### Table 8 PCR options for relative standard curve experiments

Single- or multiplex PCR	PCR or RT-PCR <sup>[1]</sup>	Detection chemistry
Singleplex	PCR	TaqMan™
Multiplex	1-step RT-PCR	SYBR™ Green
	2-step RT-PCR	

<sup>[1]</sup> RT-PCR: reverse transcription PCR.

## Comparative C<sub>t</sub> experiments

## **Overview**

Use comparative  $C_t$  ( $\Delta\Delta C_t$ ) experiments to determine relative target quantity in samples.

In a comparative C<sub>t</sub> experiment, the software performs the following tasks.

1. The software measures amplification of the target of interest and of an endogenous control target in a reference (calibrator) sample and in test samples.

The endogenous control is a target that is expressed equally in all samples; examples of endogenous controls are  $\beta$ -actin, GAPDH, and 18S ribosomal RNA. The software can algorithmically incorporate multiple endogenous control targets in relative quantification calculations.

The reference sample is used as the basis for relative quantification results (or 1× sample). For example, in a study of drug effects on gene expression, an untreated control is an appropriate reference sample.

- 2. The measurements for the target of interest are normalized to the endogenous control.
- 3. To determine the relative quantity of the target in test samples, the software compares the normalized  $\Delta C_q$  ( $\Delta C_t$  or  $\Delta C_{rt}$ ) for the sample to the normalized  $\Delta C_q$  ( $\Delta C_t$  or  $\Delta C_{rt}$ ) for the reference sample.

For a comparison of this method to the relative standard curve method, see "Relative quantitation: relative standard curve vs. comparative Ct" on page 56.

## **Reaction types**

Comparative C<sub>t</sub> experiments include the following reaction types for the endogenous control target and each target of interest.

Reaction type (task)	Sample description
Reference sample <sup>[1]</sup>	The sample that is used as the basis for relative quantification results
Unknown	Test or reference sample
No-template control (NTC/ Negative	Water or buffer
Control)	No amplification of the target should occur in NTC wells.

 Table 9
 Reaction types for comparative C<sub>t</sub> experiments

<sup>[1]</sup> To identify a sample as a reference sample, review the relative quantification settings.

The precision of quantification experiments improves as the number of replicate reactions increases. Set up the number of replicates appropriate for the experiment.



## **Compatible PCR options**

Single- or multiplex PCR	PCR or RT-PCR <sup>[1]</sup>	Detection chemistry
Singleplex	PCR	TaqMan™
Multiplex	1-step RT-PCR	SYBR™ Green
	2-step RT-PCR	

#### Table 10 PCR options for comparative C<sub>t</sub> experiments

<sup>[1]</sup> RT-PCR: reverse transcription PCR

# Relative quantitation: relative standard curve vs. comparative $\ensuremath{C_t}$

Use either relative standard curve experiments or comparative C<sub>t</sub> experiments to determine the relative quantity of a target of interest in a test sample relative to a reference sample. Relative quantitation experiments are commonly used for the following applications.

- Comparison of expression levels of a gene in different tissues.
- Comparison of expression levels of a gene in a treated sample vs. an untreated sample.
- Comparison of expression levels of a gene of interest in different genetic backgrounds.
- Analysis of the gene expression changes over time under specific treatment conditions.

Characteristic	Relative standard curve	Comparative C <sub>t</sub>	
Typical use	Best for assays that have suboptimal PCR efficiency.	Best for high-throughput measurements of relative gene expression of many genes in many samples.	
Advantage	Requires the least amount of validation because the PCR efficiencies of the target and endogenous control do not need to be equivalent.	<ul> <li>Relative levels of target in samples can be determined without the use of a standard curve, if the PCR efficiencies of the target and endogenous control are relatively equivalent.</li> <li>Reduced reagent usage.</li> <li>More space available in the reaction plate.</li> </ul>	
Limitation	A standard curve must be constructed for each target, which requires more reagents and more space in the reaction plate.	<ul> <li>Suboptimal (low PCR efficiency) assays may produce inaccurate results.</li> <li>Before you use the comparative C<sub>t</sub> method, we recommend that you determine that the PCR efficiencies for the target assay and the endogenous control assay are approximately equal.</li> </ul>	

## Set up a relative standard curve experiment in the software

- 1. In the **Home** screen, create or open a template.
  - In the 🚵 New Experiment pane, click Create New Experiment to create a new template.
  - In the 
     *Open Existing Experiment* pane, click Open to select and open an existing template.
- 2. In the **Properties** tab, enter the template information.
- In the Method tab, adjust the reaction volume.
   For most experiments, the default run method is appropriate.
- 4. In the Plate tab (Quick Setup), assign plate attributes.
  - a. In the Plate Attributes pane, select a Passive Reference, Reference Sample, and Endogenous Control from the dropdown lists.
- 5. In the **Plate** tab (**Quick Setup**), define and assign well attributes.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. Assign samples and targets to selected wells.
    - Enter new sample and target names in the text fields.
    - Select previously defined samples and targets from the dropdown lists.

**Note:** New sample or target names entered in the **Quick Setup** subtab are automatically populated with default values for **Reporter** (FAM) and **Quencher** (NFQ-MGB) and assigned a task (U **Unknown**). Edit these values in the **Advanced Setup** subtab.

- In the Plate tab, set up standard dilutions (see "Define and set up standard dilutions" on page 109).
- 7. (Optional) In the Plate tab (Advanced Setup), assign tasks.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. In the Targets table, select the checkbox of a target, then select a task from the dropdown list.

Reaction type	Task
Unknown (test sample)	U
No-template control	м

8. (*Optional*) In the **Plate** tab (**Advanced Setup**), define and assign biological replicate groups (see "Define and assign biological replicate groups" on page 110).

The targets of interest and the endogenous control target should each have wells that are assigned with standard dilutions, unknown, and no-template-control tasks, and corresponding samples.



In the Plate tab (Advanced Setup), ensure the Samples table contains the following samples.

- Unknown samples
- Reference sample
- (*Optional*) For a dilution of target standards, each dilution step for each endogenous control target and target of interest has its own sample name.

## Set up a comparative C<sub>t</sub> experiment in the software

- 1. In the **Home** screen, create or open a template.
  - In the 🚵 New Experiment pane, click Create New Experiment to create a new template.
  - In the Model Open Existing Experiment pane, click Open to select and open an existing template.
- 2. In the **Properties** tab, enter the template information.
- In the Method tab, adjust the reaction volume.
   For most experiments, the default run method is appropriate.
- 4. In the **Plate** tab (**Quick Setup**), assign plate attributes.
  - a. In the Plate Attributes pane, select a Passive Reference, Reference Sample, and Endogenous Control from the dropdown lists.
- 5. In the Plate tab (Quick Setup), define and assign well attributes.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. Assign samples and targets to selected wells.
    - Enter new sample and target names in the text fields.
    - Select previously defined samples and targets from the dropdown lists.

**Note:** New sample or target names entered in the **Quick Setup** subtab are automatically populated with default values for **Reporter** (FAM) and **Quencher** (NFQ-MGB) and assigned a task (U Unknown). Edit these values in the **Advanced Setup** subtab.

- 6. (Optional) In the Plate tab (Advanced Setup), assign tasks.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. In the **Targets** table, select the checkbox of a target, then select a task from the dropdown list.

Reaction type	Task
Unknown (test sample)	U
No-template control	м

7. (*Optional*) In the **Plate** tab (**Advanced Setup**), define and assign biological replicate groups (see "Define and assign biological replicate groups" on page 110).

The targets of interest and the endogenous control target should each have wells assigned with unknown and no-template-control tasks, and corresponding samples.

In the Plate tab (Advanced Setup), ensure the Samples table contains:

- Unknown samples
- Reference sample

## Set up and run the PCR reactions

- 1. Assemble the PCR reactions. Follow the manufacturer's instructions for the reagents and follow the plate layout set up in the software (see "Prepare reactions" on page 23).
- 2. In the desktop software, open the appropriate template (EDT file).
- **3.** Load the reaction plate into the instrument and start the run (see "Start and monitor a run" on page 24).

## **Review results**

## Workflow: Review relative standard curve and comparative $C_t$ experiments

**IMPORTANT!** If you omit wells or configure the analysis settings, click **Analyze** to reanalyze the data.

## View and assess the Standard Curve Plot

This section only applies to relative standard curve experiments.

#### Standard Curve Plot overview

The Standard Curve Plot displays the standard curve for samples designated as standards. The software calculates the quantity of an unknown target from the standard curve.

Results or metrics	Description	Criteria for evaluation
amplification calculated using the slope of the		A slope close to -3.3 indicates optimal, 100% PCR amplification efficiency.
efficiency	regression line in the standard curve.	Factors that affect amplification efficiency:
		<ul> <li>Improper design of the primer and probe</li> </ul>
		<ul> <li>Range of standard quantities—For accurate and precise efficiency measurements, use a broad range of standard quantities, 5 to 6 logs (10<sup>5</sup>- to 10<sup>6</sup>-fold).</li> </ul>
		<ul> <li>Number of standard replicates — For accurate efficiency measurements, include replicates to decrease the effects of pipetting inaccuracies.</li> </ul>
		• PCR inhibitors—PCR inhibitors and contamination in the reaction can reduce amplification efficiency.
		Other possible factors:
		<ul> <li>Component and properties of the reaction mix, such as salt content, DMSO, pH, etc.</li> </ul>
		<ul> <li>Inaccurate sample or reagent pipetting</li> </ul>
		<ul> <li>Improper analysis settings</li> </ul>
		<ul> <li>Incorrect plate setup</li> </ul>
R <sup>2</sup> value (correlation	The R <sup>2</sup> value is a measure of the closeness of fit between the	<ul> <li>A value of 1.00 indicates a perfect fit between the regression line and the data points.</li> </ul>
coefficient)	regression line and the individual $C_q$ data points of the standard reactions.	<ul> <li>An R<sup>2</sup> value &gt; 0.99 is desirable.</li> </ul>
Error	The standard error of the slope of the regression line in the standard curve.	Acceptable value is determined by the experimental criteria.
	The error can be used to calculate a confidence interval	

Table 12 Results or metrics to review in the Standard Curve Plot

(CI) for the slope and therefore the amplification efficiency.



Results or metrics	Description	Criteria for evaluation
C <sub>t</sub> values	The threshold cycle (C <sub>t</sub> ) is the PCR cycle number at which the fluorescence level meets the threshold.	<ul> <li>A C<sub>t</sub> value &gt; 8 and &lt; 35 is desirable.</li> <li>C<sub>t</sub> value &lt; 8—There may be too much template in the reaction.</li> <li>C<sub>t</sub> value &gt; 35—There may be a low amount of target in the reaction; for C<sub>t</sub> values &gt; 35, expect a higher standard deviation.</li> </ul>

Table 12 F	Results or metrics t	o review in the Standard	Curve Plot	(continued)
------------	----------------------	--------------------------	------------	-------------

#### View and assess the Standard Curve Plot

You can view and assess the Standard Curve Plot in the Results tab.

If no data are displayed in the Results tab, click Analyze.

- 1. Select Standard Curve from the dropdown list.
- 2. Click to configure the plot, then make the following selections:
  - Target: Select the target of interest
  - Plot Color: Sample, Target, or Task
  - Select all wells in the III Plate Layout

The Standard Curve Plot is displayed. The slope, R<sup>2</sup> value, amplification efficiency, and error are displayed below the plot.

- 3. Confirm that the slope, R<sup>2</sup> value, amplification efficiency, and error meet the experimental criteria.
- 4. Visually check that all unknown sample  $C_q$  ( $C_t$  or  $C_{rt}$ ) values fall within the standard curve range.
- 5. In the  $\equiv$  Well Table, use the Group By dropdown list to confirm that the C<sub>q</sub> (C<sub>t</sub> or C<sub>rt</sub>) values of all replicate samples meet the experimental criteria.

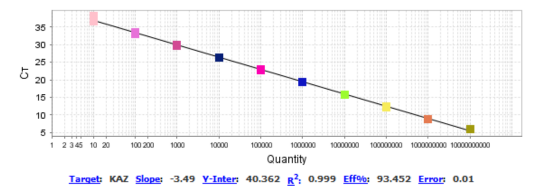


Figure 10 Example Standard Curve Plot

If the results do not meet the experimental criteria, troubleshoot using one of the following strategies:

- Omit wells, then reanalyze. For more information, see "Omit outliers from analysis" on page 35.
- Repeat the experiment, adjusting the template setup and analysis settings to improve results.

To learn more about the Standard Curve Plot, see "Standard Curve Plot overview" on page 50.

#### Standard curve settings overview

You can use the standard curve from another experiment and apply it to the current experiment. The two experiments must be from the same instrument type, block type, and run method.

To import an external standard curve, select **Analysis Settings** > Standard Curve Settings, then follow the instructions on the screen.

For step-by-step instructions for adjusting the standard curve settings, see the desktop software Help.

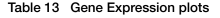
#### Gene Expression Plot overview

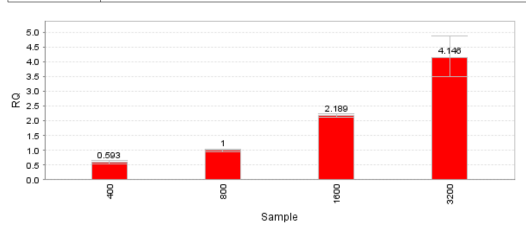
The Gene Expression Plot displays the results of relative quantification calculations for relative standard curve and comparative C<sub>t</sub> experiments.

Review the Gene Expression Plot to evaluate the fold change in expression level of the targets of interest in the test samples relative to the reference sample.

There are two plots available, depending on the experimental focus. Each plot can be viewed on a linear, log<sub>10</sub>, Ln, and log<sub>2</sub> scale.

Plot type	Description
RQ vs. Target	Groups the relative quantification (RQ) values by target. Each sample is plotted for each target.
RQ vs. Sample	Groups the relative quantification (RQ) values by sample. Each target is plotted for each sample.







In this example, there is one target of interest, and the reference sample (calibrator) is sample 800.



## QC Plot overview

The QC Plot is a visual display of the  $C_q$  ( $C_t$  or  $C_{rt}$ ) levels of potential endogenous control targets across all samples (Endogenous Control Profile).

Use the QC Plot to help choose the best endogenous control for an experiment. Select the target with a quantity (indicated by  $C_q$  ( $C_t$  or  $C_{rt}$ ) value) that does not change under experimental conditions.

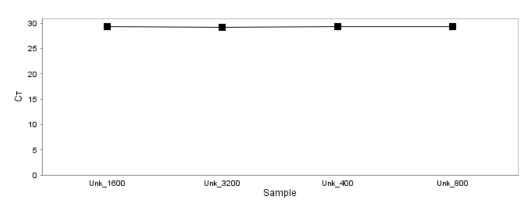
All targets can be displayed in the QC Plot. You can view up to four potential endogenous controls at a time.

#### View and assess the QC Plot

You can view and assess the QC Plot in the Results tab.

If no data are displayed, click Analyze.

- In the Results tab, select QC Plot from the dropdown list. The Endogenous Control Profile is displayed.
- 2. In the right pane, select the targets to display, then select the color and shape from the dropdown lists.
- 3. (Optional) In the View Replicate Results Table tab, select the samples to omit from analysis.
- 4. (Optional) To change the endogenous controls used for analysis, select Analysis
   Settings > Relative Quantification Settings (see "Relative quantification settings overview" on page 65).



5. Click Analyze to see the result of the adjustments.

Figure 12 Example QC Plot

## 5

## Relative quantification settings overview

In the **Results** tab, select Analysis Settings > Relative Quantification Settings to configure the following parameters:

Parameter	Description	
Analysis Type	Select Multiplex or Singleplex analysis.	
References	Set the reference sample, or set a biological replicate group as the reference sample.	
Endogenous Controls	Change the endogenous control, or select multiple endogenous controls.	
Efficiency	Set the amplification efficiency for a target.	
(Comparative C <sub>t</sub> experiments only)	The amplification efficiency for each target is calculated from the standard dilution series in relative standard curve experiments.	
Outlier Rejection (Multiplex reactions only)	Outliers with $\Delta C_q$ ( $\Delta C_t$ or $\Delta C_{rt}$ ) values less than or equal to the entered value are rejected.	
RQ Min/Max Calculations	Determines the algorithm used to calculate the relative quantification minimum and maximum values (error bars).	
	• <b>Confidence Level</b> —Select to calculate the RQ minimum and maximum values based on the selected confidence level.	
	• Standard Deviations—Select to calculate the RQ minimum and maximum values based on the selected number of standard deviations.	

For step-by-step instructions for adjusting the relative quantification settings, see the desktop software Help.



# Set up, run, and review genotyping experiments

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## **Genotyping experiments**

## Overview

Use genotyping experiments to detect single nucleotide polymorphism (SNP) variants of a target nucleic acid sequence.

Genotyping experiments use preformulated TaqMan<sup>™</sup> SNP Genotyping Assays that include the following components:

- Two sequence-specific primers for amplification of sequences containing the SNP of interest
- Two allele-specific TaqMan™ probes for Allele 1 and Allele 2

In a genotyping experiment, the software performs the following tasks.

- 1. The software normalizes the fluorescence of the reporter dyes to the fluorescence of the passive reference dye in each well.
- 2. The software plots the normalized reporter dye signal of each sample well on an Allelic Discrimination Plot, which contrasts the reporter dye intensities of the allele-specific probes.
- 3. The software algorithmically clusters the sample data, and assigns a genotype call to the samples of each cluster according to its position on the plot.

## **Reaction types**

Reaction type (task)	Sample description
Unknown	Test sample
No-template control	Water or buffer
	No amplification of the target should occur in NTC wells.
Allele control (1/1)	Control sample that is homozygous for allele 1
Allele control (1/2)	Control sample that is heterozygous allele 1/allele 2
Allele control (2/2)	Control sample that is homozygous for allele 2

Table 14 Reaction types for genotyping experiments

Allele controls are optional but recommended. Including allele controls helps to improve the clustering algorithm, particularly in situations where a limited number of samples are run.

In genotyping experiments, the software makes calls for individual wells. Running 3 or more replicates of each reaction can help identify outlier wells that may be present.

## **Compatible PCR options**

#### Table 15 PCR options for genotyping experiments

Single- or multiplex PCR	PCR or RT-PCR <sup>[1]</sup>	Detection chemistry	
Multiplex <sup>[2]</sup>	PCR	TaqMan™	

<sup>[1]</sup> RT-PCR: reverse transcription PCR

<sup>[2]</sup> Each SNP genotyping assay is a multiplex assay with a probe for each allele. Multiple SNP assays can be performed in a single well.

Genotyping calls are based either on end-point data (data collected outside of any PCR cycling stage) or on real-time data (data collected during a PCR cycling stage). For detailed information, see the analysis settings section of this guide.

We recommend collecting real-time amplification data during the PCR stage, for troubleshooting purposes.

## Set up a genotyping experiment in the software

- 1. In the **Home** screen, create or open a template.
  - In the 🚵 New Experiment pane, click Create New Experiment to create a new template.
  - In the 
     *Open Existing Experiment* pane, click Open to select and open an existing template.
- 2. In the **Properties** tab, enter the template information.
- In the Method tab, adjust the reaction volume.
   For most experiments, the default run method is appropriate.

- 4. In the Plate tab (Quick Setup), assign plate attributes.
  - a. In the Plate Attributes pane, select the Passive Reference from the dropdown list.
- 5. In the Plate tab (Quick Setup), define and assign well attributes.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. Assign samples and SNP assays to selected wells.
    - Enter new sample and SNP assay names in the text fields.
    - Select previously defined samples and SNP assays from the dropdown lists.

**Note:** New sample or SNP assay names entered in the **Quick Setup** subtab are automatically populated with the following default values:

Allele 1	Reporter	Quencher	Allele 2	Reporter	Quencher	Task
Allele 1	VIC	NFQ-MGB	Allele 2	FAM	NFQ-MGB	Unknown

Edit these values in the Advanced Setup subtab.

- 6. (Optional) In the Plate tab (Advanced Setup), assign tasks.
  - a. Select wells in the iii Plate Layout or the  $\equiv$  Well Table.
  - **b.** In the **SNP Assays** table, select the checkbox of a SNP assay, then select a task from the dropdown list.

Reaction type	Task
Unknown (test sample)	U
No-template control	И
Allele control (1/1) <sup>[1]</sup>	1
Allele control (1/2) <sup>[1]</sup>	12
Allele control (2/2) <sup>[1]</sup>	22

<sup>[1]</sup> Optional but recommended

In the Plate tab (Advanced Setup), ensure the Samples table contains the following samples:

- Unknown samples
- (Optional) Allele control samples

## Set up and run the PCR reactions

- 1. Assemble the PCR reactions. Follow the manufacturer's instructions for the reagents and follow the plate layout set up in the software (see "Prepare reactions" on page 23).
- 2. In the desktop software, open the appropriate template (EDT file).
- **3.** Load the reaction plate into the instrument and start the run (see "Start and monitor a run" on page 24).



## **Review results**

## Workflow: Review genotyping experiments Assess the Allelic Discrimination Plot (page 70) ▼ (Optional) View the Amplification Plot (page 32) ▼ Review data for outliers and (optional) omit wells (page 35) ▼ (Optional) View the Multicomponent Plot to review the dye signal profile (page 38) ▼ (Optional) View the Multicomponent Plot to review the signal profile (page 38) ▼ (Optional) View the Raw Data Plot to review the signal profile (page 39) ▼ (Optional) Review flags in the QC Summary (page 40) ▼ (Optional) Configure the analysis settings (page 71, page 115)

## Allelic Discrimination Plot overview

The Allelic Discrimination Plot contrasts the Rn or the  $\Delta$ Rn of the reporter dyes for the allele-specific probes of the SNP assay. It is an intermediary step in the software algorithm for genotyping calls.

Data points tend to cluster along the horizontal axis (Allele 1), vertical axis (Allele 2), or diagonal (Allele 1/Allele 2).

Table 16	Data clusters in the Allelic Discrimination Plot
----------	--

A substantial increase in	Clusters along	Indicates
Fluorescence of VIC <sup>™</sup> dye-labeled probe only	Horizontal axis	Homozygosity for Allele 1
Fluorescence of FAM <sup>™</sup> dye-labeled probe only	Vertical axis	Homozygosity for Allele 2
Fluorescence of both VIC™ and FAM™ dye-labeled probes	Diagonal	Heterozygosity for Allele 1 – Allele 2

Review the allelic discrimination plot to assess data clusters.

- Confirm that clustering of control samples is as expected.
- Visually assess clusters for the three possible genotypes.



**Note:** The desktop software clustering algorithm does not call genotypes if all the samples are one genotype (form one cluster).

## View and assess the Allelic Discrimination Plot

You can view and assess the Allelic Discrimination Plot in the Results tab.

If no data are displayed in the **Results** tab, click **Analyze**.

- 1. In the **Results** tab, select **Allelic Discrimination Plot** from the dropdown list.
- 2. Click ( to configure the plot, then make the following selections:
  - SNP Assay: select the assay of interest
  - Plot Type: Cartesian or Polar

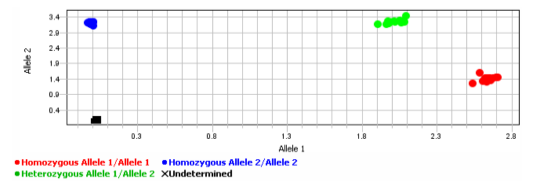
The Allelic Discrimination Plot is displayed for the selected SNP assay.

**Note:** Initially, all points in the plot are cyan because all of the wells in the **IIII Plate Layout** are selected. Click anywhere in the plot or **IIII Plate Layout** to deselect all wells. The data points in the plot change to the call colors.

- 3. Confirm that control data clusters as expected.
  - a. In the  **Well Table** or  **Plate Layout**, select the wells containing a control to highlight the corresponding data points in the plot.
  - **b.** Check that the data points for each genotype control cluster along the expected axis of the plot.
- 4. Select the cluster at the bottom-left corner of the plot, then confirm that only the negative control wells are selected in the IIII Plate Layout or ≡ Well Table.

Samples can unexpectedly cluster with the negative controls for one of the following reasons.

- Samples contain no DNA.
- Samples contain PCR inhibitors.
- Samples are homozygous for a sequence deletion.
- 5. Review the other clusters in the plot.
  - a. Click-drag a box around a cluster to select the associated wells.
  - **b.** Confirm that the expected wells are selected in the  $\blacksquare$  **Plate Layout** or  $\equiv$  **Well Table**.
- 6. Look for outliers outside the three genotype clusters.



#### Figure 13 Example Allelic Discrimination Plot

To confirm results, retest outliers and samples with no amplification (cluster with negative controls).

#### Perform manual calls

You can perform manual calls in the Results tab.

- 1. In the **Results** tab, select **Allelic Discrimination Plot** from the dropdown list.
- 2. If the data are not analyzed, click **Analyze**.
- 3. (For multiple assays only) Click ⊚, then select a SNP assay from the dropdown list.
- 4. In the Allelic Discrimination Plot, use the lasso tool to select the samples to be manually called.
- 5. Click (1), then select the allele call from the Apply Call dropdown list.
- 6. Click Analyze.

**IMPORTANT!** To maintain manual calls after reanalysis, select **Analysis Settings > Call Settings**, then deselect **Default Settings** and select **Keep Manual Calls from Previous Analysis**.

Note: To remove manual calls, select Analysis Settings > Call Settings, deselect Keep Manual Calls from Previous Analysis, then reanalyze.

#### Call settings overview (genotyping)

In the **Results** tab, select Analysis Settings > Call Settings to edit the following settings:

- Data analysis settings
- · Default call settings for SNP assays without custom call settings
- · Custom call settings for individual SNP assays



Table 17	Options for data	a analysis settings	(genotyping experiment	s)

Data analysis setting	Description
Analyze Data from Post-PCR Read Only	Only post-PCR read data is used to determine calls.
Analyze Data from Pre-PCR Read and Post-PCR Read <sup>[1]</sup>	The pre-PCR read is subtracted from the post-PCR read to determine calls.
Analyze Real-Time Rn Data <sup>[2,3]</sup>	The normalized reporter data (Rn) from the user-selected cycle of the cycling stage is used to determine calls.
Analyze Real-Time Rn - Median (Rna to Rnb) <sup>[2,4,3]</sup>	A <i>quick baseline-subtracted Rn</i> from the user-selected cycle of the cycling stage is used to determine calls. The quick baseline-subtracted Rn is the Rn minus the median value of the baseline region. The median subtraction provides improved data accuracy.
Analyze Real-Time dRn Data <sup>[2,3]</sup>	The regular $\Delta Rn$ (dRn) from the user-selected cycle of the cycling stage is used to determine calls. The $\Delta Rn$ is calculated by subtracting the best-fit line through the baseline region.
	This method is better if the baselines are not flat.

<sup>[1]</sup> The run method must include a pre-read stage.

 $\ensuremath{^{[2]}}$  Data collection must be on during the PCR stage.

<sup>[3]</sup> Analysis is not restricted to the last cycle; adjust the analysis cycle using the Reveal Traces feature while viewing the Allelic Discrimination Plot.

<sup>[4]</sup> Rna to Rnb refers to all the cycles from the Start Cycle Number to the End Cycle Number of the baseline region.

#### Table 18 Options for call settings (genotyping experiments)

Call setting	Description
Autocaller Enabled	The autocaller algorithm is used to make genotype calls.
Keep Manual Calls from Previous Analysis	If autocaller is enabled, maintains manual calls after reanalysis.
Quality Value	The Quality Value is a proprietary estimation of the likelihood that a genotyping call is correct (associated with the correct cluster). If the Quality Value is less than the setting, the call is undetermined.

For step-by-step instructions for adjusting the call settings, see the desktop software Help.



# Set up, run, and review presence/absence experiments

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# Presence/absence experiments

#### **Overview**

Use presence/absence experiments to determine the presence or absence of a target nucleic acid sequence in a sample.

The software calls the target present or absent based on an algorithmically determined call threshold. (The call threshold is different from the  $C_t$  threshold; the  $C_t$  threshold is not used to make calls.)

#### **Reaction types**

Presence/absence reaction types depend on whether the experiment is set up with or without an internal positive control (IPC).

• Presence/absence experiments with IPC (recommended) are multiplex assays for the target of interest and the IPC target. The IPC is used to confirm that a negative result for the target of interest is not caused by a failed PCR.

Table 19	Reaction types for presence/absence experiments with IPC

Reaction type (task)	Sample description
Unknown	Test sample
	and
	IPC template
Negative control	Water or buffer
	and
	IPC template
No amplification control (NAC; blocked IPC) <sup>[1]</sup>	Water or buffer plus a blocking agent
	and
	IPC template; amplification prevented by blocking agent

<sup>[1]</sup> Minimum of two replicates is required for this control.



• Presence/absence experiments without IPC are singleplex reactions.

Table 20 Reaction types for presence/absence experiments without IPC

Reaction type (task)	Sample description
Unknown	Test sample
Negative control	Water or buffer

The software makes calls for individual wells. Running three or more replicates of each reaction can help identify outlier wells that may be present.

#### **Compatible PCR options**

 Table 21
 PCR options for presence/absence experiments

Single- or multiplex PCR	PCR or RT-PCR <sup>[1]</sup>	Detection chemistry
Singleplex (without IPC)	PCR	TaqMan™
Multiplex (with IPC)	1-step RT-PCR	
	2-step RT-PCR	

<sup>[1]</sup> RT-PCR: reverse transcription-PCR

Presence/absence calls are based on end-point data (data collected after the PCR stage).

- The data collected is the normalized intensity of the reporter dye, or Rn.
- If end-point experiments include pre-PCR data points, the software calculates the delta Rn (ΔRn) value according to the following formula:

 $\Delta Rn = Rn_{(post-PCR read)} - Rn_{(pre-PCR read)}$ , where Rn = normalized readings.

We recommend collecting real-time amplification data during the PCR stage, for troubleshooting purposes.

# Set up the presence/absence experiment in the software

- 1. In the **Home** screen, create or open a template.
  - In the 🚵 New Experiment pane, click Create New Experiment to create a new template.
  - In the Some Some Source Source
- 2. In the **Properties** tab, enter the template information.
- In the Method tab, adjust the reaction volume.
   For most experiments, the default run method is appropriate.
- 4. In the **Plate** tab (**Quick Setup**), assign plate attributes.
  - a. In the Plate Attributes pane, select the Passive Reference from the dropdown list.

- 5. In the Plate tab (Quick Setup), define and assign well attributes.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. Assign samples and targets to selected wells.
    - Enter new sample and target names in the text fields.
    - Select previously defined samples and targets from the dropdown lists.

**Note:** New sample or target names entered in the **Quick Setup** subtab are automatically populated with default values for **Reporter** (FAM) and **Quencher** (NFQ-MGB) and assigned a task (U **Unknown**). Edit these values in the **Advanced Setup** subtab.

- 6. (Optional) In the Plate tab (Advanced Setup), assign tasks.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. In the Targets table, select the checkbox of a target, then select a task from the dropdown list.

Reaction type	Target	Task
Unknown (test sample)	Target of interest	U
	IPC	Ι
Negative control	Target of interest	N
	IPC	Ι
NAC (blocked IPC)	Target of interest	М
NAC (blocked if C)	IPC	×

### Set up and run the PCR reactions

- 1. Assemble the PCR reactions. Follow the manufacturer's instructions for the reagents and follow the plate layout set up in the software (see "Prepare reactions" on page 23).
- 2. In the desktop software, open the appropriate template (EDT file).
- **3.** Load the reaction plate into the instrument and start the run (see "Start and monitor a run" on page 24).

QuantStudio™ Design and Analysis Desktop Software User Guide



# **Review results**

# 

#### Presence/Absence Plot overview

The Presence/Absence Plot displays the intensity of the fluorescence for each well.

Review the Presence/Absence Plot to confirm that amplification in the control wells is as expected and to review the calls for the unknown samples.

Table 22 Expected results for control reactions

Reaction type	Target	Result	Call
Negative control	IPC	Amplification	IPC Succeeded
	Target of interest <sup>[1]</sup>	No amplification	Negative control
NAC (blocked IPC)	IPC <sup>[2]</sup>	No amplification	Blocked IPC Control
	Target of interest	No amplification	Negative control

<sup>[1]</sup> The target threshold is calculated from the negative control reactions.

<sup>[2]</sup> The IPC threshold is calculated from the NAC reactions.



#### Table 23 Criteria for calls in unknown reactions

Target signal	IPC Signal	Call
Above the target threshold	Above or below the IPC threshold	Presence
Below the target threshold	Above the IPC threshold	Absence
Below the target threshold	Below the IPC threshold	Unconfirmed

#### View and assess the Presence/Absence Plot

You can view and assess the **Presence/Absence Plot** in the **Results** tab.

If no data are displayed in the **Results** tab, click **Analyze**.

- 1. In the Results tab, select Presence/Absence Plot from the dropdown list.
- 2. Click 
  to configure the plot, making the following selections:
  - Target Reporter: target defined for the target of interest
  - Control Reporter: target defined for the IPC
  - For the initial review of the **Presence/Absence Plot**, select the following options:
    - Show Calls: All Calls
    - Show IPC
    - Show Controls

The **Presence/Absence Plot** is displayed for data points selected in the plot settings. The data points for selected wells in the  $\parallel\parallel$  **Plate Layout** or  $\equiv$  **Well Table** are highlighted in the plot (see Figure 14).

- **3.** Confirm that amplification in the negative and blocked IPC control wells is as expected. Use one of the following options:
  - Select control wells in the **IIII** Plate Layout or **E** Well Table, then confirm the location of the data points in the Presence/Absence Plot.

  - View the amplification plots for the negative controls (see Figure 15 and "Optimize display of negative controls in the Amplification Plot" on page 36).

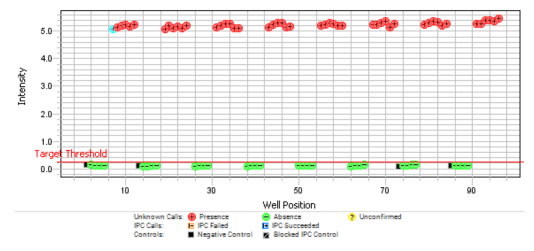
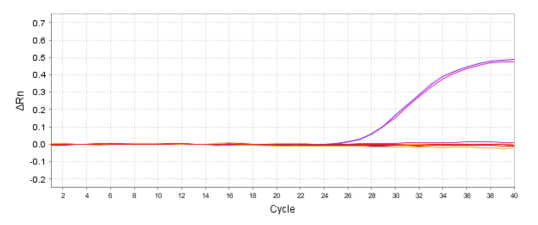
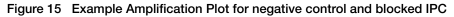


Figure 14 Example Presence/Absence Plot

The IPC results are not displayed in this example.





Amplification of the IPC target (blue lines) is seen in the negative control wells but not the blocked IPC (NAC) wells. No amplification of the target of interest (red lines) is seen in either negative control or blocked IPC wells.

#### Call settings overview (presence / absence)

Use the Call Settings tab to edit the following settings:

- Data analysis settings
- · Default call settings for assays without custom call settings
- · Custom call settings for individual assays

Table 24	Options for data	a analysis settings	(presence/absence	experiments)

Data analysis setting	Description
Analyze Data from Post-PCR Read Only	Only post-PCR read data is used to determine calls.
Analyze Data from Pre-PCR Read and Post-PCR Read	The pre-PCR read is subtracted from the post-PCR read to determine calls.

Call setting	Description
Confidence Value	<ul> <li>What confidence value is used to determine the target and IPC call thresholds.</li> <li>A lower confidence value or more controls typically results in a lower calculated threshold.</li> </ul>
	<ul> <li>threshold.</li> <li>A higher confidence value or fewer controls typically results in a higher calculated threshold.</li> </ul>

Table 25 Options for call settings (presence/absence experiments)

For step-by-step instructions for adjusting the call settings, see the desktop software Help.



# Set up, run, and review melt curve experiments

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# Melt curve experiments

#### **Overview**

Use melt curve experiments to determine the melting temperature (T<sub>m</sub>) of the amplification products of a PCR that used intercalating dyes.

Melting temperature ( $T_m$ ) is the temperature at which 50% of the DNA is double-stranded and 50% is dissociated into single-stranded DNA. The melt curve of a single amplification product displays a single peak at the product's  $T_m$ . Multiple peaks in a melt curve experiment indicate additional amplification products, usually from non-specific amplification or formation of primer-dimers.

In the software, melt curve analysis is included in the default run method for any experiment type that uses intercalating dyes.

- 1. The software plots a melt curve based on the fluorescence of the dye with respect to change in temperature.
- 2. Using the melt curve, the software calculates the melting temperature ( $T_m$ ).

#### **Reaction types**

Table 26 Reaction types for melt curve experiments

Reaction type (task)	Sample description
Unknown	Previously run PCR reactions that used intercalating dyes
No-template control (NTC/ Negative Control)	Previously run PCR reactions that used water or buffer Note: No DNA should be present in NTC wells.

# Set up a melt curve experiment in the software

- 1. In the **Home** screen, create or open a template.
  - In the 🚵 New Experiment pane, click Create New Experiment to create a new template.
  - In the Some Dependence of the Interview Depen
- 2. In the **Properties** tab, enter the template information.
- 3. In the **Method** tab, adjust the reaction volume.

(*Optional*) Edit the ramp increment in the melt curve (see "Edit the ramp increment for the melt curve dissociation step" on page 106).

- 4. In the **Plate** tab (**Quick Setup**), assign plate attributes.
  - a. In the Plate Attributes pane, select the Passive Reference from the dropdown list.
- 5. In the Plate tab (Quick Setup), define and assign well attributes.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. Assign samples and targets to selected wells.
    - Enter new sample and target names in the text fields.
    - Select previously defined samples and targets from the dropdown lists.

**Note:** New sample or target names entered in the **Quick Setup** subtab are automatically populated with default values for **Reporter** (FAM) and **Quencher** (NFQ-MGB) and assigned a task (U **Unknown**). Edit these values in the **Advanced Setup** subtab.

- 6. (Optional) In the Plate tab (Advanced Setup), assign tasks.
  - a. Select wells in the iii Plate Layout or the  $\equiv$  Well Table.
  - b. In the **Targets** table, select the checkbox of a target, then select a task from the dropdown list.

Reaction type	Task
Unknown (test sample)	U
No-template control	N

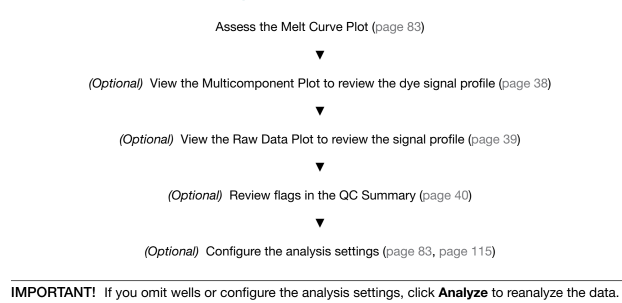


# Set up melt curve reactions

Melt curve experiments are performed using previously amplified PCR products, usually at the end of the PCR run method. You can also use a plate from an intercalating dyes-based PCR run on another instrument.

# **Review results**

#### Workflow: Review melt curve experiments



#### Melt Curve Plot overview

The Melt Curve Plot displays the melt curve of the amplification products in the selected wells.

Review the Melt Curve Plot to confirm that the amplification products in a well display a single melting temperature (T<sub>m</sub>). Multiple peaks in a melt curve indicate non-specific amplification or primer-dimer formation.

Table 27	Melt	Curve	plots
----------	------	-------	-------

Plot	Description
Derivative Reporter vs. Temperature	Displays the derivative reporter signal in the y-axis as a function of temperature. The peaks in the plot indicate significant decrease in SYBR <sup><math>m</math></sup> Green signal, and therefore the T <sub>m</sub> of the amplified products. Use this plot to confirm a single T <sub>m</sub> of the amplification products.
Normalized Reporter vs. Temperature	Displays the fluorescence signal from the reporter dye normalized to the fluorescence signal of the passive reference, as a function of temperature. You can use this plot to check the quality of the fluorescence data.

#### View and assess the Melt Curve Plot

You can view and assess the Melt Curve Plot in the Results tab.

If no data are displayed in the **Results** tab, click **Analyze**.

- 1. In the Results tab, select Melt Curve Plot from the dropdown list.
- 2. Click to configure the plot, making the following selections:
  - Plot Type: Derivative Reporter
  - Color: Sample, Target, or Well
  - Target: All or a target of interest
  - (For custom experiments with more than one Melt Curve stage) Select the Melt Curve stage to view.

The Melt Curve Plot is displayed for the selected wells of the selected stage.

- **3.** Review the plot for evidence of unexpected multiple peaks, which can indicate non-specific amplification or formation of primer-dimers.
- 4. Review the  $\equiv$  Well Table for the calculated T<sub>m</sub> in each well.

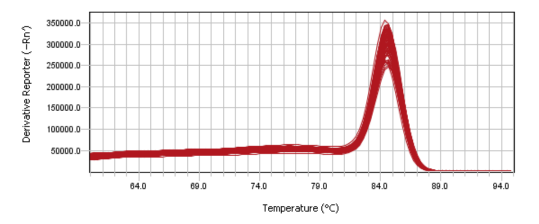


Figure 16 Example Melt Curve Plot

#### Melt curve settings overview

Use the **Melt Curve Settings** tab to enable or disable the Multi-Peak Calling function and adjust the detection levels for additional peaks, if needed.

• Enable or disable Multi-Peak Calling.

Multi-Peak Calling	Description
Enabled	<ul> <li>More than one PCR product is expected to amplify.</li> <li>T<sub>m</sub> will be determined for more than one peak.</li> </ul>
Disabled	<ul> <li>A single PCR product is expected to amplify.</li> <li>T<sub>m</sub> will be determined for one peak.</li> </ul>



• (For multi-peak calling only) Adjust the detection levels for additional peaks.

Option	Description
Peak level relative to the dominant peak (%)	Specify a fractional-level value as the additional peak detection threshold. The detected peaks are measured relative to the height of the tallest peak, which has a perfect fractional level of 100%. The default value is 10%.
	For example, set a fractional-level detection threshold value at 40, then peaks above 40% of the tallest peak are reported, and peaks below 40% are regarded as noise.
Peak Calling Threshold	Specify an absolute fluorescence-level value as the peak calling threshold. The absolute fluorescence is measured on the derivative reporter (-dRn') axis. Only peaks that appear above the peak calling threshold will be detected.
	For example, set a fluorescence-level value at 90,000, then peaks with fluorescence above 90,000 are reported, and peaks below 90,000 are regarded as noise.

For step-by-step instructions for adjusting the melt curve settings, see the desktop software Help.



# Use the software with the Security, Auditing, and E-signature (SAE) Administrator Console

## Overview of the Security, Auditing, and E-signature Administrator Console

The software is compatible with the Security, Auditing, and E-signature Administrator Console.

When you install the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, you have the option to install the SAE Administrator Console.

The information in this chapter is for the security, audit, and e-signature tasks that are performed in the QuantStudio<sup>™</sup> Design and Analysis Desktop Software.

For information about the Security, Auditing, and E-signature Administrator Console and the tasks that are performed in the console, see *SAE Administrator Console v2.1 User Guide* (Pub. No. MAN0028414).

#### Table 28 Functions of the Security, Auditing, and E-signature Administrator Console

Feature	Description
System security	Controls user access to an application. A default user account assigned the Administrator role is provided at installation. You can set up additional SAE user accounts and permissions.
Auditing	Tracks actions performed by users and changes to the SAE settings. Some actions are automatically audited silently.
	The following audit functions can be set up:
	<ul> <li>Audit changes to specific objects and specify the audit mode.</li> </ul>
	Generate reports for audited user actions and SAE function changes.
	Generate reports for software or instrument actions and runs.
Electronic signature (e-signature)	Determines if users are required to fulfill signature requirements before performing specific functions. You can perform the following functions:
	• Configure e-signature so that a user can perform an action only if the associated data are signed.
	<ul> <li>Configure each e-signature event to require one or multiple signatures and to require users with specific roles to sign.</li> </ul>



# Enable the system security function

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, click **Tools → Security → Settings**.
- 2. In the Login dialog box, enter your user name and password, then click Settings.
- 3. In the SAE Settings dialog box, select the connection option.

Option	Description	
Local server radio button	Use this option when the SAE Administrator Console and the QuantStudio™ Design and Analysis Desktop Software are installed on the same computer.	
Remote server radio button	Use this option when the SAE Administrator Console and the QuantStudio™ Design and Analysis Desktop Software are installed on separate computers.	

4. If a remote server was selected, enter the IP address of the computer that is running the SAE Administrator Console in the **Server IP address** field.

Port number 8443 must be used. Do not edit this value. Ensure that this port is open on the network.

Depending on the way that the network is configured, the IP address of the computer might change. This affects the connection between the SAE Administrator Console and the QuantStudio<sup>™</sup> Design and Analysis Desktop Software. A DHCP network connection with a reserved IP address is recommended. A static IP address can also be used.

- 5. Click Log In.
- 6. In the SAE Settings dialog box, select the Enable SAE settings checkbox, then click Save.
- 7. Click OK.

# Disable the system security function

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, click **Tools > Security > Settings**.
- 2. In the SAE Settings dialog box, deselect the Enable SAE settings checkbox, then click Save.
- 3. Click **Yes** to confirm.
- 4. Click **OK** to confirm.

Files that were created or edited with the security function enabled can be opened and edited after the security function is disabled. The audit and e-signature information for these files is not available after the security function is disabled. Any edits that are made to the files do not have any audit information collected. E-signatures are not collected.

The audit and e-signature information that was collected before the security function was disabled is retained. It can be accessed if the security function is enabled again.

# **Overview of accounts**

You must sign in to the software when security is enabled. A user name and a password are required.

Accounts are set up and edited in the Security, Auditing, and E-signature Administrator Console.

Accounts are associated with permissions for specific tasks in the software.

For more information about setting up and editing accounts, see *SAE Administrator Console v2.1 User Guide* (Pub. No. MAN0028414).

In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, individual users can change their own password. Administrators can change passwords for any account in the Security, Auditing, and E-signature Administrator Console.

Depending on how the account is set up in the Security, Auditing, and E-signature Administrator Console, an individual user might need to reset their password the first time they log in to the QuantStudio<sup>™</sup> Design and Analysis Desktop Software.

# Log in to QuantStudio<sup>™</sup> Design and Analysis Desktop Software

If security is enabled in the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, the **Login** dialog box is displayed when the software is launched.

Ensure that the administrator has set up an account in the Security, Auditing, and E-signature Administrator Console.

- 1. In the Login dialog box, enter your user name in the User Name field, then enter your password in the Password field.
- 2. Click Log In.

Depending on how the account is set up in the Security, Auditing, and E-signature Administrator Console, an individual user might need to reset their password the first time they log in to the QuantStudio<sup>™</sup> Design and Analysis Desktop Software.

# Change your password in QuantStudio<sup>™</sup> Design and Analysis Desktop Software

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, click **Tools → Security → Change Password**.
- 2. Enter your password in the Old Password field.
- 3. Enter your new password in the New Password field and the Confirm Password field.
- 4. Click OK.



# Log out of QuantStudio<sup>™</sup> Design and Analysis Desktop Software

In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, click **Tools → Security → Log Out**.

# Default roles and permissions

**IMPORTANT!** Permissions apply to all of the user accounts that are assigned to the role.

Table 29	Default roles and	permissions
Table Lo	Bondant ronoo ana	pormooreme

Permissions		Roles		
Category	Function	Administrator	Scientist	Technician
Setup	Create new template	Yes	Yes	Yes
	Edit template properties	Yes	Yes	Yes
	Edit run method	Yes	Yes	No
	Edit analysis settings	Yes	Yes	No
	Omit wells for analysis	Yes	Yes	No
	Define new target/assay	Yes	Yes	No
	Define new sample	Yes	Yes	Yes
	Edit sample	Yes	Yes	Yes
	Delete sample	Yes	Yes	Yes
	Assign targets/assays	Yes	Yes	Yes
	Assign samples	Yes	Yes	Yes
	Edit passive reference	Yes	Yes	Yes
Run	Start a run from the desktop software	Yes	Yes	Yes
	Stop a run from the desktop software	Yes	Yes	Yes
Target library	Create targets	Yes	Yes	Yes
	Edit targets	Yes	Yes	Yes
	Delete targets	Yes	Yes	No
Analysis settings library	Create analysis settings	Yes	Yes	Yes
	Edit analysis settings	Yes	Yes	Yes
	Delete analysis settings	Yes	Yes	No

Permissions		Roles		
Category	Function	Administrator	Scientist	Technician
Dye library	Create a custom dye	Yes	Yes	Yes
	Delete a custom dye	Yes	Yes	No
Preferences	Edit system preferences	Yes	Yes	No
Edit file save destination	Edit save as destination	Yes	Yes	No
	Edit export destination	Yes	Yes	No
	Edit report destination	Yes	Yes	No
Security configuration	Log into timed-out user sessions	Yes	No	No
	Perform e-signing	Yes	Yes	Yes
Security configuration (Security,	Configure security and auditing	Yes	No	No
Auditing, and E-signature Administrator Console)	View action records	Yes	No	No
	View system configuration records	Yes	No	No
	View application object records	Yes	No	No
	View instrument run records	Yes	No	No

Table 29 Default roles and permissions (continued)

# View and manage the audit records

#### View the object audit records

The object audit records are maintained within the template file or the data file. The audit records are not displayed in the SAE Administrator Console.

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, open a template file or a data file.
- 2. Select the Audit tab.

The audit records for the file are displayed. There are two panes, a **Summary** pane and a **Details** pane. The **Summary** pane lists the audit records. Each audit record corresponds to an update of the file. The **Details** pane lists each individual edit to a file.

- 3. In the **Summary** pane, select a row that corresponds to an audit record. The details for the specific audit record are displayed in the **Details** pane.
- 4. (Optional) Click Z Action > Show Filter to display the filter pane at the top of the screen. If the filter pane is displayed, the button is **Hide Filter**.

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5. Select the Filter by checkbox to filter the audit records.

The audit records can be filtered by the following items:

- Date range
- User name
- Full name
- Record type
- Record name
- Reason
- Action

**IMPORTANT!** When entering a value in a field, the text must match the exact text in the audit record. It is case-sensitive and the punctuation must match. If the text does not match the audit record, no results are displayed.

6. Click Refresh to update the results.

The results display the audit records that correspond to the filters. The audit records that correspond to the filters are displayed in the **Summary** pane. Select a row to view the details in the **Details** pane.

Some of the filters are based on the details. For example, if you filter by an action, all of the audit records that contain this action in the details are displayed in the **Summary** pane.

#### Generate an object audit report

All of the audit records are included in the report. This applies even if filters were applied in the software.

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, open a template file or a data file.
- 2. Select the Audit tab.
- Click Action > View Report.
   The Print Preview dialog box is opened and it displays a preview of the audit report.
- 4. Click Save As PDF, select a folder location, enter a file name, then click Save.
- 5. Click Print Report to print the file.

#### Export the audit summary

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, open a template file or a data file.
- 2. Select the Audit tab.
- 3. In the **Summary** tab, select a row that corresponds to an audit record.
- 4. Click Action > Export Summary.
- 5. In the **Save** dialog box, select a folder location, enter a file name, then click **Save**.

The summary is exported in a CSV file format.

#### Export the audit details

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, open a template file or a data file.
- 2. Select the Audit tab.
- 3. In the **Summary** tab, select a row that corresponds to an audit record.
- 4. Click Z Action > Export Details.
- 5. In the Save dialog box, select a folder location, enter a file name, then click Save.

The details are exported in a CSV file format. The information that is included in the file is from the **Details** pane that is associated with the selected audit record.

# View and manage the e-signatures

#### Sign the data

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, open a template file or a data file.
- 2. Click Tools > Security > Sign data.
- **3.** In the **Electronic Signature** dialog box, select a meaning from the **Meaning** dropdown list. The meanings that are available in the dropdown lists correspond to the meanings that were set up in the Security, Auditing, and E-signature Administrator Console.
- 4. Enter your user name and password, then click Sign Data.

The electronic signature is available in the **e-sig** tab.

#### View the e-signatures

- 1. In the QuantStudio<sup>™</sup> Design and Analysis Desktop Software, open a template file or a data file.
- 2. Select the e-Sig tab.

All of the e-signatures for the file are displayed.

- 3. (Optional) Select a row to view a preview of the experiment report in the bottom pane.
- 4. (Optional) Save or print the experiment report.
  - Click **Save As PDF**, select a folder location, enter a file name, then click **Save**.
  - Click **Print Report**, then select the print settings.
- 5. (Optional) Click ☑ Action → Print e-Signatures.

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- 6. (Optional) In the **Print Preview** dialog box, save or print the e-signature history report.
  - Click Save As PDF, select a folder location, enter a file name, then click Save.
  - Click **Print Report**, then select the print settings.



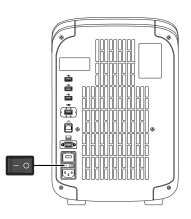
# Instrument overview

Power on the instrument	93
View run history and delete or transfer files from the instrument	93
Load and unload the plate in the instrument	94

### Power on the instrument

- 1. Touch anywhere on the touchscreen to determine if the instrument is in sleep mode. If the home screen is displayed, the instrument is already powered on.
- 2. If the home screen does not display, power on the instrument by pressing the switch on the rear panel.

If left unattended (for about two hours), the instrument automatically enters sleep mode (enabled by default) to conserve power.



# View run history and delete or transfer files from the instrument

In the home screen, touch ( Settings > Run History.

- Touch an individual run record to view its details, then complete one of the following actions:
  - Touch **Delete** to delete the run record.
  - Touch **Transfer** to export the run data.
- Touch Manage to select multiple run records for simultaneous viewing, deletion, or transfer.



#### Note:

- Guests (users not signed-in) can only view guest run records.
- Users signed into their instrument profiles can also view their own run records.
- · Administrators can view all run records.

**Note:** If the connection between the instrument and the desktop software is interrupted during the run, the instrument still completes the run. However, the run data (EDS file) must be transferred from the instrument to the desktop software using a USB drive or a network drive.

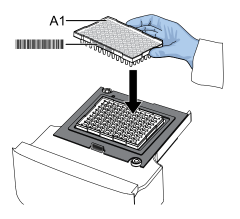
# Load and unload the plate in the instrument

#### Load and unload a plate in the QuantStudio<sup>™</sup> 1 Real-Time PCR Instrument



CAUTION! Use flat caps for 0.2-mL tubes. Rounded caps can damage the heated cover.

- 1. Load the plate.
  - a. Open the instrument drawer.
  - b. Load the plate onto the plate adapter so that the following criteria are met.
    - Well A1 of the plate is in the top-left corner of the plate adapter.
    - The barcode faces the front of the instrument.



**Note:** Do not remove the black plate adapter before loading a plate or tube strips. If used, tube strips can fit loosely in the adapter, but the heated cover will apply the appropriate pressure to seat the tube strips securely in the adapter.

- c. Close the instrument drawer.
- 2. When the run ends, unload the plate.
  - a. Open the instrument drawer.
  - b. Remove the plate.



c. Close the instrument drawer.



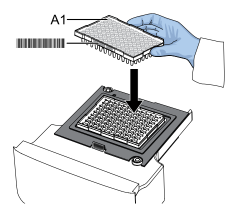
**CAUTION! PHYSICAL INJURY HAZARD.** During instrument operation, the plate temperature can reach 100°C. Allow it to cool to room temperature before handling.

# Load and unload a plate in the QuantStudio<sup>™</sup> 3 Real-Time PCR Instrument or QuantStudio<sup>™</sup> 5 Real-Time PCR Instrument



**CAUTION!** Use flat caps for 0.2-mL tubes and 0.1-mL tubes. Rounded caps can damage the heated cover.

- 1. Load the plate.
  - a. Touch (a) to eject the instrument drawer.
  - **b.** Load the plate onto the plate adaptor so that the following criteria are met.
    - Position well A1 of the plate in the top-left corner of the plate adapter.
    - Ensure that the barcode faces the front of the instrument.



**IMPORTANT!** The instrument should be used by trained operators who have been warned of the moving parts hazard.

**Note:** (*For 96-well 0.2-mL blocks only*) Do not remove the black plate adapter before loading a plate or tube strips. If used, tube strips can fit loosely in the adapter, but the heated cover will apply the appropriate pressure to seat the tube strips securely in the adapter.

**Note:** The 384-well and 96-well Fast (0.1-mL) block configurations do not require a plate adapter.

- c. Touch a to close the instrument drawer.
- 2. When the run ends, unload the plate.
  - a. Touch (a) to eject the instrument drawer.
  - b. Remove the plate.



**c.** Touch (a) to close the instrument drawer.



**CAUTION! PHYSICAL INJURY HAZARD.** During instrument operation, the plate temperature can reach 100°C. Allow it to cool to room temperature before handling.

Note: If the instrument does not eject the plate, contact Support.



# Alternative procedures to set up a template

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Assign targets, samples, and biological replicate groups from an XLS file	101
Create new EDT files using existing EDT and EDS files	102

## Set up a custom experiment

#### Custom experiments overview

Custom experiment setup is required for assays that use multiple PCR stages, such as TaqMan<sup>™</sup> Mutation Detection Assays. A custom experiment also allows flexibility for secondary analysis.

The default settings for custom experiments are that of a standard curve experiment, but most settings are editable.

Setting	Default
Run method (thermal protocol)	Equivalent to standard curve experiment default
Tasks	Unknown 🕕
	Negative control
	Standard S
C <sub>t</sub> settings	Baseline threshold equivalent to standard curve experiment default
Flag settings	QC flags on
	No automatic omissions
Auto Export	Off

Table 30 Default settings in custom experiments
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Table 31	Editing options in custom experiments
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Setting	Description
Run method (thermal protocol)	<ul> <li>Ramp rates can be edited within software limits.</li> <li>Data collection can be enabled at any step and during any ramp within a melt stage.</li> <li>Multiple instances of any type of stage can be added, with exceptions noted.</li> <li>Any stage can be added at any point in a run method, with exceptions noted.</li> <li>Noted exceptions: <ul> <li>Only one infinite hold (must be added at the end).</li> <li>Only one pre-PCR read and one post-PCR read stage. If both exist in a run method, the pre-read must be before the post-read.</li> </ul> </li> <li>For example, the following order is valid: melt-PCR-Pre-Read-Melt-PCR.</li> </ul>
Analysis settings	Editable
Flags	Editable
Optical filters	Editable

#### Set up a custom experiment in the software

- 1. In the **Home** screen, create or open a template.
  - In the 🚵 New Experiment pane, click Create New Experiment to create a new template.
  - In the Some Some Source Source
- 2. In the Properties tab, enter the template information.
- 3. In the Method tab, edit the default run method according to the experiment requirements.
- 4. In the Plate tab (Quick Setup), assign plate attributes.
  - a. In the Plate Attributes pane, select the Passive Reference from the dropdown list.
- 5. In the Plate tab (Quick Setup), define and assign well attributes.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - b. Assign samples and targets to selected wells.
    - Enter new sample and target names in the text fields.
    - Select previously defined samples and targets from the dropdown lists.

**Note:** New sample or target names entered in the **Quick Setup** subtab are automatically populated with default values for **Reporter** (FAM) and **Quencher** (NFQ-MGB) and assigned a task (U **Unknown**). Edit these values in the **Advanced Setup** subtab.

- 6. (Optional) In the Plate tab (Advanced Setup), assign tasks.
  - a. Select wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
  - **b.** In the **Targets** table, select the checkbox of a target, then select a task from the dropdown list.
- 7. (*Optional*) In the **Plate** tab (**Advanced Setup**), define and assign biological replicate groups (see "Define and assign biological replicate groups" on page 110).

# Assign samples using a sample definition file

#### About sample definition files

Import sample information from a sample definition file to include in the plate setup. A sample definition file is a CSV file or a TXT file that contains the following setup information:

- Sample name
- (Optional) Custom sample properties

#### Create a sample definition file

- 1. In a spreadsheet program, create the following column headers:
  - Well
  - Sample Name
  - (*Optional*) Column header names for up to 32 user-defined custom fields (for example, **Custom 1**, **Custom 2**, etc.)
- 2. Enter the well number and sample name in the appropriate columns.
- 3. (Optional) Enter the custom properties for the sample.
- 4. Save the file as a tab-delimited text file (TXT) or a comma-separated values file (CSV).

#### Import sample information from a sample definition file

Example setup files are provided with the software in:

<drive>:\Program Files (x86)\Applied Biosystems\QuantStudio Design and Analysis Software\examples\User Sample Files,

where *<drive>* is the drive on which the software is installed.

- 1. In an open experiment, select File > Import Plate Setup.
- 2. Click Browse, navigate to a sample definition text file, then click Select.
- 3. Click Apply.
- 4. If the experiment already contains plate setup information, the software prompts for the replacement of the plate setup with the data from the file. Click **Yes** to replace the plate setup information.

The samples appear in the **Samples** table for the experiment. All samples and well assignments in the experiment are replaced with those in the file. If defined, the custom sample properties also appear in the  $\equiv$  **Well Table** of the **Results** tab and in the  $\equiv$  **Plate Layout** tooltips in both the **Plate** and **Results** tabs. The custom fields can be exported with the results data.

**Note:** To modify custom sample properties information, edit the custom fields in the sample definition file and import the file again. All of the sample information in the experiment is replaced with the information in the new file.

### Assign samples and targets using plate setup files

#### About plate setup files

Plate setup files contain setup information such as the well number, sample name, sample color, target name, dyes, and other reaction plate contents.

Plate setup files can be exported from previously run experiments. For instructions on exporting an experiment, see "Export experiments or results" on page 42.

#### Import plate setup data

Import the plate setup for a new experiment from an exported file with one of the following formats:

- EDS EDS file format
- EDT user-created and system templates files format
- TXT text format
- XML XML format
- CSV comma separated values format
- SDT Sequence Detection System (SDS) template files format
- SDS 7900 v2.4 format

**Note:** Import plate setup information from a 96-well plate into a 384-well plate, provided that the sample file is a TXT file.

**IMPORTANT!** The file must contain only plate setup data and it must match the experiment type.

- 1. In the Plate tab, select File > Import Plate Setup.
- 2. Click **Browse**, navigate to and select the file to import, then click **Select**.

Example setup files are provided with the software in: <drive> :\Program Files (x86) \Applied Biosystems\QuantStudio Design and Analysis Software\examples\User Sample Files,

- where <drive> is the drive on which the software is installed.
- 3. Click Apply.

The setup data from the selected file is imported into the open experiment.

# Assign targets, samples, and biological replicate groups from an XLS file

For wells with single targets, you can paste assignment information from an XLS file into the plate layout of the desktop software.

An example copy and paste file is provided with the software in:

```
<drive>:\Program Files (x86)\Applied Biosystems\QuantStudio Design and
Analysis Software\examples\User Sample Files.
```

where *<drive>* is the drive on which the software is installed.

- 1. In the custom properties tab of the example Microsoft<sup>™</sup> Excel<sup>™</sup> file, ensure that the **Well** column is sorted in order 1 through 96, then select the **Well** column and the **Sample Name** column, *including* headers.
- 2. In the **Plate** tab of the software, click  $\equiv$  **Well Table**, then ensure that the well numbers are in order from 1 through 96.
- In the ≡ Well Table, hover the mouse in the first cell underneath the Sample header (adjacent to A1), right-click, then select either Paste or Paste only samples.
   Any of the columns not copied are treated as NULL values for those columns.

QuantStudio™ Design and Analysis Desktop Software User Guide

# Create new EDT files using existing EDT and EDS files

#### About experiment templates

Use templates to create experiments with the same parameters or with pre-existing settings. Experiments can be saved as unlocked or locked (password-protected) templates.

You can save the following information in an experiment template (EDT) file:

- Plate setup information (defined sample and targets or SNP assays, plate assignment of samples and targets or SNP assays)
- Reagent information
- Run method (thermal protocol)
- Analysis settings

Example templates are provided with the software in:

<drive>:\Program Files (x86)\Applied Biosystems\QuantStudio Design & Analysis
Software\templates,

where *<drive>* is the drive on which the software is installed.



# Detailed procedures to create or edit a method

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# Adjust method parameters

For an overview of the method as it is graphically represented, see "Method elements" on page 19.

- In the **Method** tab, click a method parameter field to edit the following information:
  - Reaction volume
  - Temperature ramp rate
  - Step temperature
  - Step hold time
  - Number of cycles
- Click-drag  $\equiv$  to increase or decrease a step temperature.
- Click o to switch data collection on or off at each step.

**Data Collection On** enables analysis of data that is collected throughout the PCR, for real-time analysis and troubleshooting.

 (QuantStudio<sup>™</sup> 3 Real-Time PCR Instrument and QuantStudio<sup>™</sup> 5 Real-Time PCR Instrument only) Click to configure settings for Auto Delta or VeriFlex<sup>™</sup> Zones for individual steps (see "Set up advanced temperature zones (Auto Delta and VeriFlex<sup>™</sup> Zones)" on page 104).

Note: In melt curve stages, Advanced Settings are not applicable.

An A or V is displayed in the PCR stage when **Auto Delta** or VeriFlex<sup>M</sup>, respectively, is enabled.

- Click III to configure pause settings.
- Adjust the heated cover temperature via the instrument settings (see the instrument user guide).



- See the Help to learn more about adjusting the following parameters:
  - Adding or subtracting a stage
  - Adding or subtracting a step from a stage
  - Configuring optical filter settings

# Set up advanced temperature zones (Auto Delta and VeriFlex<sup>™</sup> Zones)

(QuantStudio<sup>™</sup> 3 Real-Time PCR System and QuantStudio<sup>™</sup> 5 Real-Time PCR System only) Configure settings for **Auto Delta** and **VeriFlex<sup>™</sup> Zones**.

- Auto Delta—Incremental increase or decrease of a cycle's temperature or hold time for a step in a cycling stage (not applicable for Hold or Infinite Hold stages).
- VeriFlex<sup>™</sup> Zones Independent temperature zones within 5°C of adjacent zones.
  - QuantStudio<sup>™</sup> 3 Real-Time PCR Instrument: 3 zones
  - QuantStudio<sup>™</sup> 5 Real-Time PCR Instrument: 6 zones

Note: VeriFlex<sup>™</sup> Zones temperature settings are not available for 384-well blocks.

1. In the **Method** tab, click 🔅 **Advanced Settings** in a step.

Note: Any changes apply only to the step in which you clicked.

- 2. Configure either the VeriFlex<sup>™</sup> Zones or Auto Delta for the selected step.
  - Select **VeriFlex**<sup>™</sup>, then enter a temperature for each zone.

Note: In the Plate tab, the VeriFlex<sup>™</sup> Zones display on the plate layout.

- To view setting details, hover over the  ${\bf V}$  in each zone.
- To hide the display of zones, click *I* Action → Hide VeriFlex<sup>™</sup> Zones.
- Select **Auto Delta**, then enter a starting cycle, temperature, and time.
- 3. Click Save.

### Add or adjust a pause step

- 1. In the **Method** tab, click <u>iii</u> in the step.
- 2. Select Pause.
- 3. Enter the cycle after which the pause should occur.
- 4. Enter a pause temperature between 4°C and 99.9°C.



**CAUTION! PHYSICAL INJURY HAZARD.** During instrument operation, the plate temperature can reach 100°C. If you want to access the plate during a run pause, enter room temperature as the pause temperature and allow the plate to cool to room temperature before handling.

- 5. Click outside of the pause dialog box to return to the method.
- 6. (Optional) To remove a pause, click . , then deselect Pause.

### Select optical filters

The need to edit optical filter settings is rare, and it is intended for advanced or custom uses only. For information on the dyes read by each filter, see the instrument user guide.

Use the optical filters settings feature to select a filter set to match the profile of a custom dye.

1. In the Method tab, select 🗹 Action > Optical filter settings.

The excitation (x) and emission (m) wavelengths that correspond to each filter are shown on the screen.

2. Select the check boxes to enable or disable filters.

A **Melt Curve Filter** table is accessible if the method contains a melt curve stage. Otherwise, use the **PCR Filter** table to select optical filters.

- 3. (Optional) Click Revert to Defaults to reset filters.
- 4. Click Close.

# Edit the ramp increment for the melt curve dissociation step

In the **Method** tab, you can perform the following tasks to edit the ramp increment for the melt cure dissociation step.

• Select the ramp increment method for the dissociation step (located under the graphical representation of the thermal protocol).

Option	Description
Continuous (default)	Continuously increases the temperature by the ramp increment (°C/sec).
Step and Hold	Increases the temperature by the ramp increment (°C), then holds at that temperature for the specified time.
No. of Data Points per Degree	Increases the temperature by the ramp increment (°C) and collects the specified number of data points per degree increased.

- (For all options) Edit the temperature ramp increment.
  - a. Click the ramp increment element in the **Dissociation** step.
  - b. Enter a value or use the up/down arrows (default is 0.15°C/s).
- (Step and Hold only) Edit the hold time after each temperature increase.
  - a. Click the time field next to Step and Hold.
  - b. Enter a value or use the up/down arrows (default is 5 seconds).
- (*No. of Data Points per Degree only*) Edit the number of data points to be collected with each degree increase.
  - a. Click the number of data points element in the **Dissociation** step.
  - b. Enter a value or use the up/down arrows (default is 10 data points).



# Detailed procedures to set up plate / well details and libraries

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# Assign well attributes (Quick Setup subtab)

In the **Quick Setup** subtab, assign well attributes by direct entry into text fields or by selecting userdefined samples and targets or SNP assays from dropdown lists.

- 1. In the Plate tab, click Quick Setup.
- 2. Select plate wells in the  $\blacksquare$  Plate Layout or the  $\equiv$  Well Table.
- 3. Assign the well attributes for the selected wells.
  - Into the text fields, enter the names of the new sample and the new target or SNP assay.
  - From the dropdown lists, select a user-defined sample and target or SNP assay. For more information about defining or importing samples and targets or SNP assays, see "Define and assign well attributes (Advanced Setup subtab)".

**Note:** In the **Advanced Setup** subtab, change the default selections for the reporter and quencher dyes and for tasks (see "Assign a task to wells" on page 108).

4. (Optional) Enter comments for the selected wells.

# Define and assign well attributes (Advanced Setup subtab)

In the **Advanced Setup** subtab, define or import samples and targets or SNP assays, then assign well attributes.

- 1. In the Plate tab, click Advanced Setup.
- 2. In the Samples table, define samples (see "Define samples in the Samples table" on page 112).

- 3. In the Targets or SNP Assays table, define targets or SNP assays, then select detection tasks.
  - **a.** Define targets or SNP assays (see "Define targets in the Targets table" on page 113 or "Define SNP assays in the SNP Assays table" on page 114, respectively).
  - b. Select a detection task from the Task column dropdown list (see "Assign a task to wells" on page 108).
- 4. Assign well attributes.
  - a. Select plate wells in the IIII Plate Layout or the ≡ Well Table (see "Select plate wells" on page 21).
  - **b.** Select the checkbox of a defined sample.
  - c. Select the checkbox of a defined target or SNP assay.

## Assign a task to wells

- 1. In the Plate tab, click Advanced Setup.
- 2. Select plate wells in the IIII Plate Layout or the ≡ Well Table (see "Select plate wells" on page 21).
- 3. In the Targets or SNPs table, select the check box of a target or SNP assay.
- 4. Select a detection task from the Task column dropdown list.

#### Detection tasks for targets and SNP assays

Task	Description
Unknown (default)	The well contains test samples with unknown genotype.
Negative Control / No template control	The well contains water or buffer instead of sample.
Standard <sup>[1]</sup>	The well contains samples with known standard quantities.
	<b>Note:</b> For a standard detection task, enter the standard quantity in the quantity column.
Positive Control Allele 1 / Allele 1 <sup>[2]</sup>	The well contains samples homozygous for allele 1.
<sup>2</sup> 2Positive Control Allele 2 / Allele 2 <sup>[2]</sup>	The well contains samples homozygous for allele 2.
Positive Control Allele 1 / Allele 2 <sup>[2]</sup>	The well contains samples heterozygous for allele 1 and 2.

(continued)

Task	Description
Internal positive control <sup>[3]</sup>	The PCR reaction contains a short synthetic DNA template to distinguish between true negative results (that is, the target is absent in the samples) and negative results caused by PCR inhibitors, incorrect assay setup, or reagent or instrument failure.
Blocked IPC <sup>[3]</sup>	The well contains an IPC blocking agent, which blocks amplification of the IPC.
NAC – No amplification control <sup>[3]</sup>	The PCR reaction contains an IPC blocking agent instead of sample. No amplification should occur in negative control-blocked IPC wells because the reaction contains no sample and amplification of the IPC is blocked.

<sup>[1]</sup> For standard curve and relative standard curve experiments only.

<sup>[2]</sup> For genotyping experiments only.

<sup>[3]</sup> For presence/absence experiments only.

## Define and set up standard dilutions

Note: This information is applicable for standard curve and relative standard curve experiments only.

- 1. In the Plate tab, select if Action > Define and Set Up Standards.
- 2. Select Singleplex or Multiplex from the Model dropdown list.
- 3. (Optional) Select the target from the dropdown list.
- 4. Enter the parameters for the dilution series.
  - Number of dilution points-5 recommended
  - Number of replicates 3 recommended
  - Starting Quantity—The highest or lowest standard quantity, without units.

Note: The quantity can be expressed as copies, copies/ $\mu$ L, ng/ $\mu$ L, or as relative dilutions.

**Note:** Use **E** to indicate the exponent number using scientific notation. For example, to indicate  $1.23 \times 10^4$ , enter 1.23E4.

Serial Factor

Note: The serial factor calculates quantities for all standard curve points.

- Starting quantity is the highest value—Select 1:10 to 1:2.
- Starting quantity is the lowest value—Select 2× to 10×.

**Note:** The **Standard Curve Preview** y-axis values are calculated from the starting quantity and serial factor. Actual results may differ from the preview.

QuantStudio™ Design and Analysis Desktop Software User Guide



- 5. Select and arrange the wells to use for the standards.
  - Select Automatically Select Wells for Me.
  - Select Let Me Select Wells, then select wells using the displayed plate layout.
- 6. Select to arrange the standards in Columns or Rows.
- 7. (Optional) Click Reset to revert to default values.
- 8. Click Apply, then click Close to return to the Plate tab.

#### Assign the standard dilutions manually

Note: Applicable for standard curve and relative standard curve experiments only.

- 1. In the Plate tab, select wells in the  $\blacksquare$  Plate Layout or  $\equiv$  Well Table.
- 2. Select the check box for the target, select S from the Task dropdown list, then enter a quantity.
- 3. Repeat to complete the standard dilution series.

## Define and assign biological replicate groups

Biological Replicate Groups can be used in standard curve, relative standard curve, comparative C<sub>t</sub>, and custom experiments.

- 1. In the Plate tab, click Advanced Setup.
- 2. Define Biological Replicate Groups.
  - a. In the Biological Replicate Groups table, click + Add.
  - b. (Optional) Click a cell to edit color, name, or comments.
  - c. (Optional) Click  $\times$  to delete a biological replicate group from the table.
- 3. Assign Biological Replicate Groups.

  - b. In the Biological Replicate Groups table, select the check box of a biological replicate group.

## Sample, target, and SNP assay libraries

#### Libraries overview

Libraries contain saved information to reuse in future templates.

The following libraries are available in the software:

- Dye Library
- Sample Library
- Target Library
- SNP Assay Library
- Analysis Settings Library

To access the libraries, use the Tools menu or the Plate tab.

- In the menu bar, select Tools > (Library of choice).
- In the Plate tab of an open EDT or EDS file, click Advanced Setup, then select ☑ Action ➤ Import from Library.

#### Permissions for the libraries

If security is enabled in the software, the following permissions are applicable:

- Targets library
  - Create Targets / Assays
  - Edit Targets / Assays
  - Delete Targets / Assays
- Analysis settings library
  - Create Analysis Settings
  - Edit Analysis Settings
  - Delete Analysis Settings
- Dye library
  - Add New Custom Dye
  - Delete Custom Dye

#### Apply a filter to search a library

You can filter the Sample, SNP assay, Target, and Analysis Settings Libraries.

- 1. Access a library of interest.
  - In the menu bar, select Tools > (Library of interest).
  - In the Plate tab of an open EDT or EDS file, click Advanced Setup, then select
     ☑ Action ➤ Import from Library.
- 2. Select a feature from the first dropdown list. Each column of the table is an available feature.



- **3.** Select a condition from the second dropdown list to define the feature. The conditions will vary by feature.
- 4. Enter a value or text by which to filter.
- 5. Click Apply Filter.

#### Define samples in the Samples table

In the **Plate** tab, click **Advanced Setup**, then perform one of the following actions in the **Samples** table.

Option	Action	
Manually define a sample	<ul> <li>a. Click + Add.</li> <li>b. Click a cell to edit the attributes for the sample.</li> <li>c. (<i>Optional</i>) Click + in the table header to add a Custom Attribute column.</li> <li>a. Click the Custom Attribute column header, then edit the header with a new sample attribute.</li> <li>b. Click a Custom Attribute cell in the table, then enter the attribute information.</li> </ul>	
Import samples from a TXT or XLS file	<ul> <li>a. Select  Action ▶ Import from File.</li> <li>b. Navigate to and select a file, then click Open.</li> </ul>	
Import samples from the Sample Library	<ul> <li>a. Select  Action → Import from Library.</li> <li>b. (Optional) Apply a filter to search for a specific sample (see page 111).</li> <li>c. Select one or more samples, then click Add Selected.</li> <li>Note: Shift-click or Ctrl-click to select multiple samples.</li> </ul>	
Save a sample to the Sample Library	Select a sample row, then select <i>I</i> Action ► Save to Library.	
Delete a sample from the table	Select a sample row, then click $\mathbf{X}$ .	

## Define targets in the Targets table

In the Plate tab, click Advanced Setup, then perform one of the following actions in the Targets table.

Option	Action
Manually define a target	a. Click + Add.
	b. Click a cell to edit the attributes for the target.
Import targets from a	a. Select <b>I</b> Action ▶ Import from Library.
TXT or XML file	b. Click Import or Import AIF.
	c. Navigate to and select a file, then click <b>Import</b> .
	d. Select one or more targets, then click <b>Add Selected</b> .
	Note: Shift-click or Ctrl-click to select multiple targets.
Import targets from the <b>Target</b>	a. Select <b>⊘ Action → Import from Library</b> .
Library	b. (Optional) Apply a filter to search for a specific target (see page 111).
	c. Select one or more targets, then click <b>Add Selected</b> .
	Note: Shift-click or Ctrl-click to select multiple targets.
Save a target to the Target Library	Select a target row, then select <i>I</i> Action ► Save to Library.
Delete a target from the table	Select a target row, then click $\mathbf{X}$ .



## Define SNP assays in the SNP Assays table

In the **Plate** tab, click **Advanced Setup**, then perform one of the following actions in the **SNP Assays** table.

Option	Action	
Manually define a SNP assay	<ul><li>a. Click + Add.</li><li>b. Click a cell to edit the attributes for the SNP assay.</li></ul>	
Import SNP assays from a TXT or XML file	<ul> <li>a. Select  Action &gt; Import from Library.</li> <li>b. Click Import or Import AIF.</li> <li>c. Navigate to and select a file, then click Import.</li> <li>d. Select one or more SNP assays, then click Add Selected.</li> </ul> Note: Shift-click or Ctrl-click to select multiple SNP assays.	
Import SNP assays from the SNP Assay Library	<ul> <li>a. Select  Action &gt; Import from Library.</li> <li>b. (<i>Optional</i>) Apply a filter to search for a specific SNP assay (see page 111).</li> <li>c. Select one or more SNP assays, then click Add Selected.</li> <li>Note: Shift-click or Ctrl-click to select multiple SNP assays.</li> </ul>	
Save a SNP assay to the SNP Assay Library	Select a SNP assay row, then Action > Save to Library.	
Delete a SNP assay from the table	Select a SNP assay row, then click $\mathbf{X}$ .	



# Configure analysis settings

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This section describes the analysis settings that apply to all experiment types, unless otherwise noted.

## Guidelines for the analysis settings

- We recommend analyzing the experiment with the default analysis settings.
- If the default analysis settings are not suitable for the experiment, modify the settings in the **Analysis Settings** dialog box, then reanalyze the experiment.
- Save modified analysis settings to the Analysis Settings Library.

The default analysis settings are different for each experiment type. The analysis settings determine for following parameters.

- How the baseline, threshold, and threshold cycle (Ct) are calculated
- Which flags are enabled
- · Other analysis options that are specific to an experiment type

For detailed information about different types of analysis settings, see the following sections.

- "Ct settings overview" on page 116
- "Flag settings overview" on page 117
- "Advanced settings overview" on page 117
- "Standard curve settings overview" on page 52
- "Relative quantification settings overview" on page 65
- "Call settings overview (genotyping)" on page 71
- "Call settings overview (presence / absence)" on page 78
- "Melt curve settings overview" on page 83

## View and configure the analysis settings

If security is enabled in the software, your account must have the permission of **Edit Analysis Settings** to configure the settings.

- 1. In the top-right corner of the **Results** tab, click .
- View and (*optional*) configure the analysis settings.
   For step-by-step instructions for adjusting analysis settings, see the desktop software Help.
- 3. Click Apply.
- 4. Click Analyze to reanalyze to experiment with the new settings.
- 5. (Optional) To save the settings in the Analysis Settings Library, click Save.
- 6. (Optional) To return to the default settings, click Revert.

## Ct settings overview

The default C<sub>t</sub> settings are appropriate for most applications. Configuration of the settings is an option for analysis of atypical or unexpected run data.

For step-by-step instructions for adjusting the C<sub>t</sub> settings, see the desktop software Help.

Note: The C<sub>t</sub> Settings feature is not available for experiments without a PCR stage, such as melt curve experiments.

Table 32	Ct Settings
----------	-------------

Setting	Description
Data Step Selection	Determines the stage/step combination for $C_t$ analysis (when there is more than one data collection point in the run method).
Algorithm Settings – Baseline Threshold	The <b>Baseline Threshold</b> Algorithm is used to calculate the $C_t$ values. This algorithm is an expression estimation algorithm that subtracts a baseline component and sets a fluorescence threshold in the exponential region.
Algorithm Settings – Relative Threshold	The Relative Threshold ( $C_{rt}$ ) Algorithm is used to calculate the $C_{rt}$ values.
Default C <sub>t</sub> Settings	Determines how the <b>Baseline Threshold</b> Algorithm is set. The <b>Default C<sub>t</sub> Settings</b> are used for targets unless they have custom settings. For recommendations on adjusting baseline and threshold settings, see Table 33.

Table 32 Ct Settings (continued)

Setting	Description
C <sub>t</sub> Settings for Target	• Default Settings selected—The Default C <sub>t</sub> Settings are used to calculate the C <sub>t</sub> values for the target.
	• <b>Default Settings</b> deselected—The software allows manual setting of the baseline or the threshold.
	For recommendations for adjusting baseline and threshold settings, see Table 33.

#### Table 33 Recommendations for manual threshold and baseline settings

Setting	Recommendation	
Threshold	Enter a value for the threshold so that the threshold is:	
	Above the background.	
	<ul> <li>Below the plateau and linear phases of the amplification curve.</li> </ul>	
	Within the exponential phase of the amplification curve.	
Baseline	Select the <b>Start Cycle</b> and <b>End Cycle</b> values so that the baseline ends before significant fluorescence signal is detected.	

## Flag settings overview

Use the Flag Settings to configure the following parameters.

- Adjust the sensitivity so that more wells or fewer wells are flagged.
- Change the flags that are applied by the software for each experiment type.

For step-by-step instructions for configuring the flag settings, see the desktop software Help.

## Advanced settings overview

Use the Advanced Settings tab to change baseline settings for individual wells.

For step-by-step instructions for adjusting the advanced settings, see the desktop software Help.

**Note:** The **Advanced Settings** feature is not available for experiments without a PCR stage, such as melt curve experiments.



# Related documentation for the QuantStudio<sup>™</sup> 1 Real-Time PCR System

Document	Publication number
QuantStudio™ 1 Real-Time PCR System Installation, Use, and Maintenance Guide	MAN0017853
QuantStudio™ Design and Analysis Desktop Software Command-Line Application Guide	MAN0010409
QuantStudio™ 1 Real-Time PCR System Site Preparation Guide	MAN0017854

# Related documentation for the QuantStudio<sup>™</sup> 3 Real-Time PCR System and QuantStudio<sup>™</sup> 5 Real-Time PCR System

Document	Publication number
QuantStudio™ 3 and 5 Real-Time PCR Systems Installation, Use, and Maintenance Guide	MAN0010407
QuantStudio™ 5 Real-Time PCR System Installation, Use, and Maintenance Guide (for China)	MAN0028412
QuantStudio™ Design and Analysis Desktop Software Command-Line Application Guide	MAN0010409
SAE Administrator Console v2.1 User Guide	MAN0028414
QuantStudio™ 3 and 5 Real-Time PCR Systems Site Preparation Guide	MAN0010405

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## Glossary

#### biological replicates

Reactions that contain identical components and volumes, but evaluate separate samples of the same biological source (for example, samples from three different mice of the same strain, or separate extractions of the same cell line or tissue sample).

For runs that use biological replicate groups in a gene expression project, the values displayed in the biological replicates lists are calculated by combining the results of the separate biological samples and treating this collection as a single population (that is, as one sample).

For  $\Delta C_t$  computations (normalizing by the endogenous control) in a singleplex experiment, the software averages technical replicates. The averages from the technical replicates are then averaged together to determine the value for that biological replicate.

#### endogenous control

A gene that is used to normalize template differences and sample-to-sample or run-to-run variation.

#### endpoint read

See post-PCR read.

#### post-PCR read

In genotyping and presence/absence experiments, the part of the instrument run that occurs after amplification. In genotyping experiments, fluorescence data collected during the post-PCR read are displayed in the allelic discrimination plot and used to make allele calls. In presence/absence experiments, fluorescence data collected during the post-PCR read are displayed in the presence/absence plot and used to make detection calls. Also called endpoint read.

#### pre-PCR read

In genotyping and presence/absence experiments, the part of the instrument run that occurs before amplification. The pre-PCR read is optional but recommended. Fluorescence data collected during the pre-PCR read can be used to normalize fluorescence data collected during the post-PCR read.

#### reference sample

In relative standard curve and comparative  $C_t$  ( $\Delta\Delta C_t$ ) experiments, the sample used as the basis for relative quantification results. Also called the calibrator.

#### target

The nucleic acid sequence that is amplified and detected during PCR.

#### task

In the software, the type of reaction performed in the well for the target.

#### technical replicates

Reactions that contain identical components and volumes, and that evaluate the same sample; important for evaluating precision.

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