QX200 Droplet Reader and QX Developer Software
User Guide

Version 1.1
Bio-Rad Technical Support Department

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Chapter 1 Safety and Regulatory Compliance

This section cites regulatory requirements for laboratory and electrical equipment, as well as requirements for working with chemicals and hazardous substances, and also explains safety precautions and recommendations.

Regulatory Compliance

The QX200 Droplet Reader has been tested and found to be in compliance with all applicable requirements of the following safety and electromagnetic standards:

- IEC 61010-2-081:2015, EN 61010-2-081:2015. Safety requirements for electrical equipment for measurement, control, and laboratory use. Part 2-081: Particular requirements for automatic and semi-automatic laboratory equipment for analysis and other purposes (includes Amendment 1)
- IEC 61010-2-101:2015 (2nd ed.). Safety requirements for electrical equipment for measurement, control, and laboratory use. Particular requirements for in vitro diagnostic (IVD) medical equipment
- IEC 61326-1:2012 (Class A), EN 61326-1:2013 (Class A). Electrical equipment for measurement, control, and laboratory use. EMC requirements, Part 1: General requirements
- CAN/CSA 22.2 No 61010-1-04, Safety requirements for electrical, equipment for measurement, control, and laboratory use, Part 1: General requirements
- Restriction of hazardous substances (ROHS) directive (European Union)
- Registration, evaluation, authorization and restriction of chemicals (REACH). European Chemicals Agency (ECHA) June 1, 2007
- Waste electrical and electronic equipment (WEEE) directive

This equipment has been tested and found to comply with the limits for a Class A digital device pursuant to Part 15 of FCC rules. These limits are designed to provide reasonable protection against harmful interference when the equipment is operated in a commercial environment.

The CE mark indicates that the manufacturer ensures the product conforms with the essential requirements of the applicable EN directives.
Chapter 1 Safety and Regulatory Compliance

The SA mark indicates that a project has been tested to Canadian and U.S. standards, and it meets the requirements of those applicable standards.

The Waste Electrical and Electronic Equipment (WEEE) Directive symbol indicates that when the end-user wishes to discard this product, it must be sent to separate collection facilities for recovery and recycling.

Safety Warning Labels

Warning labels posted on the instrument and in this manual warn you about sources of injury or harm. Table 1 defines each safety warning label.

Table 1. Meaning of safety warning labels

<table>
<thead>
<tr>
<th>Icon</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Warning Icon]</td>
<td><strong>Warning about risk of harm to body or equipment</strong>&lt;br&gt;Operating the QX200 Droplet Reader before reading this manual can constitute a personal injury hazard. For safe use, do not operate this instrument in any manner unspecified in this manual. Only qualified laboratory personnel trained in the safe use of electrical equipment should operate this instrument. Always handle all components of the system with care and with clean, dry hands.</td>
</tr>
<tr>
<td>![Biohazard Icon]</td>
<td><strong>Warning about handling biohazardous materials</strong>&lt;br&gt;When handling biohazardous samples, adhere to the recommended precautions and guidelines and comply with any local guidelines specific to your laboratory and location.</td>
</tr>
</tbody>
</table>
Safe Use Specifications

For safe operation of the QX200 Droplet Reader, Bio-Rad Laboratories, Inc. strongly recommends that you comply with instructions listed in this section and in Instrument Maintenance on page 187.

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

- This instrument is for use only by trained personnel.
- Use only the power cord supplied with the instrument, and the plug adapter corresponding to the electrical outlets in your region.
- Position the instrument on a solid, stable surface, with adequate room at the back and on each side so that users can easily reach the power cord and USB port.
- This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the provided instructional documentation, may cause harmful interference to radio communications. Operation of the systems in a residential area is likely to cause harmful interference, in which case users will be required to correct the interference at their own expense.

**Note:** Bio-Rad recommends maintaining a backup power source in case of power outages.

Table 2 lists the safe use specifications for Bio-Rad’s QX200 Droplet Reader. The supplied shielded cables must be used with these instruments to ensure compliance with the Class A FCC limits.

**Table 2. Conditions for safe use**

<table>
<thead>
<tr>
<th>Usage aspect</th>
<th>Conditions for safe use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rated input power</td>
<td>Input: 100–240 V, 50–60 Hz, 1A</td>
</tr>
<tr>
<td></td>
<td>Voltage fluctuations shall not exceed +10% of ratings for the included external power supply.</td>
</tr>
<tr>
<td></td>
<td>Use only the power cord supplied with the equipment.</td>
</tr>
<tr>
<td></td>
<td>Fuse: 10A 250V SLOW BLOW</td>
</tr>
<tr>
<td>Pollution degree/environment</td>
<td>2 (indoor use only)</td>
</tr>
<tr>
<td>Usage temperature</td>
<td>18–30°C</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>85% maximum (non-condensing)</td>
</tr>
</tbody>
</table>
### Usage aspect

<table>
<thead>
<tr>
<th>Usage aspect</th>
<th>Conditions for safe use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Altitude</td>
<td>0 to 6,560 ft (0 to 2,000 meters) above sea level</td>
</tr>
<tr>
<td>Installation category</td>
<td>II (external power supply plugs into a standard AC receptacle)</td>
</tr>
<tr>
<td>Ventilation requirements</td>
<td>The following distances should be unobstructed for proper ventilation:</td>
</tr>
<tr>
<td></td>
<td>■ 5 in (12 cm) on the left and right sides of the instrument</td>
</tr>
<tr>
<td></td>
<td>■ 10 in (25 cm) behind the machine</td>
</tr>
<tr>
<td>Benchtop requirements</td>
<td>The following requirements apply to the benchtop or table on which the instrument will be installed.</td>
</tr>
<tr>
<td></td>
<td>The benchtop must be 76&quot; W x 30&quot; D (193cm x 76cm) single benchtop rated for more than 250 lbs (114 kg) load</td>
</tr>
<tr>
<td></td>
<td><strong>Important:</strong> Do not place instrument over multiple benchtop surfaces.</td>
</tr>
<tr>
<td></td>
<td>34&quot; – 40&quot; (86cm – 102cm); actual height is at the discretion of user for an ergonomic operation of the equipment</td>
</tr>
</tbody>
</table>

### Hazards

The QX200 Droplet Reader is designed to operate safely when used in the manner prescribed by the manufacturer. If the instrument or any of its associated components is used in a manner not specified by the manufacturer, the inherent protection provided by the instrument may be impaired.

Bio-Rad Laboratories, Inc. is not liable for any injury or damage caused by the use of this equipment in any unspecified manner, or by modifications to the instrument not performed by Bio-Rad or an authorized agent. Only trained Bio-Rad personnel should perform service on the QX200 Droplet Reader.

### Biohazards

The QX200 Droplet Reader is a laboratory product. However, if biohazardous samples are present, adhere to the following guidelines and comply with any local guidelines specific to your laboratory and location.

**Note:** No biohazardous substances are expended during normal operations of this instrument.
General Precautions

- Always wear laboratory coat, laboratory gloves, and safety glasses with side shields or goggles.
- Keep your hands away from your mouth, nose, and eyes.
- Completely protect any cut or abrasion before working with potentially infectious materials.
- Wash your hands thoroughly with soap and water after working with any potentially infectious material before leaving the laboratory.
- Store all infectious or potentially infectious material in unbreakable leak-proof containers.
- Before leaving the laboratory, remove protective clothing.
- Do not use a gloved hand to write, answer the telephone, turn on a light switch, or touch anything that other people may touch without gloves.
- Change gloves frequently. Remove gloves immediately when they are visibly contaminated.
- Do not expose materials that cannot be properly decontaminated to potentially infectious material.
- Upon completion of an operation involving biohazardous material, decontaminate the work area with an appropriate disinfectant (for example, a 1:10 dilution of household bleach).

Surface Decontamination

**WARNING!** To prevent electrical shock, always turn off and unplug the instrument prior to performing decontamination procedures.

**Important:** Do not use abrasive or corrosive detergents or strong alkaline solutions. These agents can scratch surfaces and damage the system.

The following areas can be cleaned with 10% bleach solution:

- Outer area and chassis
- Inner plate holders
- Droplet reading surfaces

To prepare and apply the disinfectant, refer to the instructions provided by the product manufacturer. For more information on surface cleaning, see Appendix C, Instrument Maintenance. For questions regarding the use of other cleaning agents, contact Bio-Rad Technical Support.

**Important:** Do not clean the handler Y-axis rail when the front door is open. This is a lubricated surface, and failures will occur if the lubrication is removed.
Disposal of Biohazardous Material

Dispose of the following potentially contaminated materials in accordance with laboratory local, regional, and national regulations:

- Clinical samples
- Reagents
- Used reaction vessels or other consumables that may be contaminated

Chemical Hazards

The QX200 Droplet Reader contains no potentially hazardous chemical materials.

Explosive or Flammability Hazards

The QX200 Droplet Reader poses no uncommon hazard related to flammability or explosion when used in a proper manner as specified by Bio-Rad.

Electrical Hazards

The QX200 Droplet Reader poses no uncommon electrical hazard to operators if installed and operated properly without physical modification and connected to a power source of proper specification.
Decommissioning and Disposal

The QX200 Droplet Reader contains electrical materials that should be disposed of as unsorted waste and must be collected separately, according to European Union Directive 2012/19/EU on waste electrical and electronic equipment — WEEE Directive. The purpose of decommissioning is to make sure that the equipment is electrically and environmentally safe for disposal. Before disposal, contact your local Bio-Rad representative for country-specific instructions.

Transport

Before moving or shipping the QX200 Droplet Reader, decontamination procedures must be performed. Always move or ship the instrument with the supplied packaging materials, which will protect the instrument from damage. If appropriate containers cannot be found, contact your local Bio-Rad office.

Warranty

The QX200 Droplet Reader and its associated accessories are covered by a standard Bio-Rad warranty. Contact your local Bio-Rad office for the details of the warranty.

Use of unapproved supermixes may harm the instrument, and voids the warranty. Alteration of this instrument voids the warranty and safety certification, as it creates a potential safety hazard.
Chapter 1 Safety and Regulatory Compliance
Chapter 2  Introduction to Droplet Digital PCR

Droplet digital polymerase chain reaction (ddPCR) is a digital PCR method based on water-oil emulsion droplet technology. ddPCR uses a combination of microfluidics and proprietary surfactant chemistries to divide each sample into water-in-oil droplets. The technology uses reagents and workflows similar to those used for most standard TaqMan probe-based assays, and provides absolute quantification of nucleic acid target sequences by counting nucleic acid molecules encapsulated in discrete, volumetrically defined water-in-oil droplet partitions.

ddPCR is highly effective in the following areas:

- **Absolute quantification** — ddPCR provides a concentration of target DNA copies per input sample without the need for running standard curves, making this technique ideal for target DNA measurements, viral load analysis, and microbial quantification.

- **Genomic alterations such as gene copy number variation (CNV)** — CNVs result in too few or too many dosage-sensitive genes responsible for phenotypic variability, complex behavioral traits, and disease. ddPCR enables measurement of 1.2x differences in gene copy number.

- **Detection of rare sequences** — researchers must amplify single genes in a complex sample, such as a few tumor cells in a wild-type background. ddPCR is sensitive enough to detect rare mutations or sequences.

- **Gene expression and microRNA analysis** — ddPCR provides stand-alone absolute quantification of expression levels, especially low-abundance microRNAs, with sensitivity and precision.

- **Next-generation sequencing (NGS)** — ddPCR quantifies NGS sample library preparations to increase sequencing accuracy and reduce run repeats. Validate sequencing results such as single nucleotide polymorphisms or copy number variations with absolute quantification.

- **Single cell analysis** — the high degree (10-fold to 100-fold) of cell-cell variation in gene expression and genomic content among homogeneous post-mitotic, progenitor, and stem cell populations drives a need for analysis from single cells. ddPCR enables low copy number quantification.

- **Genome edit detection** — ddPCR enables fast, precise, and cost-effective assessment of HDR (Homology directed repair) and NHEJ (non-homologous end joining) generated by CRISPR-Cas9 or other genome editing tools.
ddPCR has the following benefits for nucleic acid quantification:

- **Unparalleled precision** — The massive sample partitioning afforded by ddPCR enables small fold differences in target DNA sequence between samples to be reliably measured.

- **Increased signal-to-noise ratio** — High-copy templates and background are diluted, effectively enriching template concentration in target-positive partitions. This allows for the sensitive detection of rare targets and enables a ±10% precision in quantification.

- **Removal of PCR efficiency bias** — Error rates are reduced by removing the amplification efficiency reliance of qPCR, enabling accurate quantification of targets.

- **Simplified quantification** — There is no requirement for a standard curve for absolute quantification.

**ddPCR Workflow**

The ddPCR process adheres to the following workflow:

- You prepare your samples for PCR by combining DNA or RNA with primers, probes dye, and Bio-Rad ddPCR supermix.

- A droplet generator fractionates a sample into approximately 20,000 uniform nanoliter-sized droplets, with target and background DNA distributed randomly into the droplets during the partitioning process.

- A thermal cycler performs PCR amplification of the nucleic acid target in each individual droplet.

- A droplet reader reads each droplet to determine the fraction of positive droplets in the original sample. Positive droplets containing at least one copy of the target DNA molecule exhibit increased fluorescence compared to negative droplets.
QX200 Droplet Digital PCR System

As part of its QX200 Droplet Digital PCR System, Bio-Rad provides the following instruments:

- QX200 Droplet Generator or the Automated Droplet Generator to generate the droplets from your samples
- QX200 Droplet Reader to read the generated and thermal-cycled droplets

Important: Your QX200 Droplet Reader performs the post-PCR droplet reading function only. A droplet generator and a thermal cycler are required to perform a full ddPCR cycle. You can purchase additional instruments, including a thermal cycler, as well as associated accessories and consumables, from Bio-Rad. For information, see Appendix G, Ordering Information.

Finding Out More

Click the Help tab and then click the Bio-Rad Website link to access links to technical notes, manuals, videos, product information, and technical support. The website also provides many technical resources on a wide variety of methods and applications related to PCR, droplet digital PCR, and gene expression.
Chapter 2  Introduction to Droplet Digital PCR
Chapter 3 About the QX200 Droplet Reader

Following the droplet generation and thermal cycling processes, the QX200 Droplet Reader collects binary data for analysis as it

- Singulates the droplets in each sample
- Streams the droplets in single-file past a two-color detector to determine which droplets contain one or more target molecules (positive reading) and which do not (negative reading)

Use the information in this chapter to correctly assemble your plate, connect the instrument, start up the instrument, and insert the plate for processing. Information on components and specifications is also included.

LEGEND

1. Button to open and close lid
2. Status indicator lights (for information, see Status Indicator Lights on page 25)
3. On the back of the instrument, power and USB outlets
4. Oil and waste compartment (for information, see Fluid Maintenance on page 187)
Included Items and Additional Components

This section identifies the functional items included with your droplet reader purchase, as well as additional components and consumables. For information on items you may need for a full ddPCR cycle, see Appendix G, Ordering Information.

Table 3 explains the components that ship with the QX200 Droplet Reader.

Table 3. QX200 Droplet Reader Included Components

<table>
<thead>
<tr>
<th>Component</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>QX200 Droplet Reader</td>
<td>Droplet reading instrument, part of QX200 ddPCR system</td>
<td>1864003</td>
</tr>
<tr>
<td>USB cable and power cord</td>
<td>USB cable that connects the computer to the instrument and power cord that connects the instrument to the power source</td>
<td>Included</td>
</tr>
</tbody>
</table>

*Note:* To obtain cables or power cords, contact Bio-Rad technical support.

Droplet reader plate holders (2) | Positions and secures the 96-well plate in the droplet reader plate compartment | 12006834        

To ensure you have everything you need to complete runs on the QX200 Droplet Reader and analyze your results, Table 4 identifies additional components and accessories that you can purchase separately.

Table 4. Additional Accessories and Consumables

<table>
<thead>
<tr>
<th>Item</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Computer</td>
<td>Computer that connects to the QX200 Droplet Reader for data collection and analysis</td>
<td>17006483</td>
</tr>
<tr>
<td>Software purchase</td>
<td>Bio-Rad QX Developer software</td>
<td></td>
</tr>
<tr>
<td>96-well PCR plates</td>
<td>ddPCR 96-well plates</td>
<td>12001925</td>
</tr>
<tr>
<td>Rainin pipets</td>
<td>20 μl for sample loading 50 μl for droplet transfer 8-channel, 200 μl for oil</td>
<td>L-20, L8-20, L-50, L8-50, L8-200</td>
</tr>
</tbody>
</table>
### Table 4. Additional Accessories and Consumables, continued

<table>
<thead>
<tr>
<th>Item</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rainin pipet tips</td>
<td>Filtered</td>
<td>GP-L10F</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GP-L200F</td>
</tr>
<tr>
<td>Foil plate seals</td>
<td>Pierceable foil plate seals</td>
<td>1814040</td>
</tr>
<tr>
<td>Plate sealer</td>
<td>PX1 PCR plate sealer</td>
<td>1814000</td>
</tr>
<tr>
<td>Droplet reader oil</td>
<td>ddPCR droplet reader oil</td>
<td>1863004</td>
</tr>
</tbody>
</table>

**Note:** Use the empty oil bottle as your waste container bottle.

### Instrument Specifications

This section contains information on the instrument size and weight of the instrument. Environmental and safe use information is explained in Safe Use Specifications on page 11.

<table>
<thead>
<tr>
<th>Element</th>
<th>Specifications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight</td>
<td>56.6 lbs (26 kg)</td>
</tr>
<tr>
<td>Size (W x H x D)</td>
<td>26 x 20.5 x 11.5” (660 x 521 x 292 cm)</td>
</tr>
</tbody>
</table>
Assembling a Plate for the QX200 Droplet Reader

The QX200 Droplet Reader accepts a solid 96-well plate that fits into a secure holder. Three separate components comprise the QX200 Droplet Reader plate:

- A base section to hold the plate
- The plate itself
- A top section to secure the plate

You must assemble the plate components before you can insert the plate into the QX200 Droplet Reader.

To assemble the plate

1. Place the plate into the base section.
2. Raise the release tabs on the top section.
3. Place the top section over the plate, and then lower the release tabs to secure the plate in the holder.
Connecting and Starting the Instrument

To execute runs on the QX200 Droplet Reader, you must connect the instrument to a computer on which the QX Developer software is installed and configured.

**Tip:** Bio-Rad recommends purchasing at least one configured computer that can remain connected to the QX200 Droplet Reader and be used for adding and running plates in QX Developer. For information on additional items you can purchase for your QX200 Droplet Reader, see Included Items and Additional Components on page 22.

**To connect the computer to the instrument**

1. Connect the USB cable provided by Bio-Rad to the USB port on the back of the QX200 Droplet Reader, and then to the computer.
2. Press the Power button on the QX200 Droplet Reader.
   The Power indicator light turns solid green to indicate that power is on.
3. Power up and log into the connected computer, and then open QX Developer and sign in.
   For information, see Signing Into the Software on page 31.

**Note:** Users must sign into the computer and software with the same secure Windows account.

After recognizing the connected instrument, QX Developer displays the Add Plate window.

**Note:** If the software does not recognize a connection to the instrument, displays the Data Analysis window.

**Status Indicator Lights**

This section describes the indicator lights on the front of the QX200 Droplet Reader.
Before you start a run, check the indicator lights to ensure the instrument is ready for droplet reading runs. For information, see Table 6.

**Table 6. Droplet Reader Status Indicator Lights**

<table>
<thead>
<tr>
<th>Indicator</th>
<th>(Power)</th>
<th>(Fluids)</th>
<th>(Plate)</th>
<th>(Run Status)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid Green</td>
<td>Power on</td>
<td>Fluid levels OK</td>
<td>Plate loaded properly</td>
<td>Run complete</td>
</tr>
<tr>
<td></td>
<td></td>
<td>See Note (1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flashing Green</td>
<td>N/A</td>
<td>Less than 30% oil is</td>
<td>N/A</td>
<td>Run in progress</td>
</tr>
<tr>
<td></td>
<td></td>
<td>available</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>More than 70% waste in</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>container</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>See Note (2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flashing Amber</td>
<td>N/A</td>
<td>Less than 10% oil is</td>
<td>N/A</td>
<td>Error during run</td>
</tr>
<tr>
<td></td>
<td></td>
<td>available</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>More than 90% (700ml) waste in container</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>See Note (3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No color</td>
<td>Power off</td>
<td>N/A</td>
<td>No plate loaded</td>
<td>Instrument is idle</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**

(1) If the bottle indicator light is solid green, there is enough oil in the bottle to complete a run on 96 wells. You can configure the full plate in QX Developer for your run.

(2) If the bottle indicator light is flashing green, there is enough oil in the bottle to start a run on a small number of wells. For example, you can run 24 wells with the oil bottle at a 19% level. If you configure your plate for more wells than the oil level allows, or if you fail to exclude the proper number of wells, the software does not enable the Start Run button.

(3) You must add oil and remove waste before the instrument can start the run.
Inserting an Assembled Plate into the Instrument

To insert a plate into the QX200 Droplet Reader

1. Press the button to open the compartment.

2. Fit the assembled plate into the compartment. Well A1 must be in the top-left position.

   Important: Make sure the tabs on the plate holder are securely clamped and the plate lays flat. Otherwise, the sampling needle can strike the lid and damage the instrument.

3. Press the button again to close the lid.

For information on adding your plate and starting runs in QX Developer, see Running Experiments on page 41.
Chapter 3 About the QX200 Droplet Reader
Chapter 4 About QX Developer

QX Developer, when connected to your QX200 Droplet Reader provides all necessary functionality to create, run, and analyze experiments on your samples.

QX Developer provides features to permit operation in compliance with Title 21 of the U.S. Code of Federal Regulations Part 11 (21 CRF Part 11) within a closed system. A closed system is defined as “an environment in which system access is controlled by the persons who are responsible for the content of electronic records that are on the system” (Section 11.3(b) (4)).

Important:

- The security controls built into QX Developer must be properly configured and administered by the software administrators in your organization in order to be secure and in compliance with 21 CFR Part 11.

- Bio-Rad makes no claim that QX Developer is CFR-compliant in and of itself, nor does the company guarantee compliance for the user. Your organization must establish policies and standard operating procedures that work in conjunction with the tools provided by Bio-Rad to ensure compliance with 21 CFR Part 11.

Using the instrument and software, you can

- Set up customized, real-time ddPCR experiments for plate runs
- Use the live analysis function during the droplet reading process
- Analyze your data files in a variety of charts and tables in the Analysis module
- Reprocess data with updated color calibration values
- Create and store plate and report templates, and produce reports on your data
- View system and experiment audit logs
- Produce an audit log report on old and new data values after reprocessing a file

Tip: If QX Developer goes into standby mode after being idle, tap or click in the lockout screen to log into the software again and redisplay the last active window.
Chapter 4 About QX Developer

The following image highlights functional areas.

1. The status bar displays information about the instrument and user. For information on specific functionality, see Instrument Status Bar on page 37.

2. Tabs provide access to the main functional windows.

3. The main pane displays the details of the selected tab.

LEGEND
Chapter 5 Getting Started in QX Developer

Use the information in this chapter to

- Sign into the software or change users
- Understand your assigned user privileges and modify your personal preferences
- Become familiar with the functional windows
- Understand the compatible file types for your software edition

Signing Into the Software

Your system administrator sets up users and privileges for the software, and communicates the information to you.

**Important:** In the Regulatory Edition, you are prompted to sign in again before the software can execute an auditable action.

**To sign in**

1. To open the software, right-click the software icon on the Desktop.
Chapter 5 Getting Started in QX Developer

The Sign in dialog box opens.

2. Enter your user name.
3. If the Sign in to: label shows the correct domain name, enter only the user name. If the label is blank or shows a different domain then enter it as domain name\user name

   **Note:** The domain and computer names shown in the graphics are examples only. If you are unsure of the domain name, contact your system administrator.

4. Enter the password, and then tap or click Sign in.

   On first use, you must agree to the end user license agreement.

5. Clear the Show EULA on sign in checkbox, and then tap or click I Agree.

   **Note:** If you tap or click I Disagree, the application closes immediately. If you leave the checkbox selected, you must agree to the EULA each time you log into the application.

   When you agree to the EULA, the dialog box closes and the application opens. Your user name appears in the Instrument Status bar.

   If you are connected to the QX200 Droplet Reader, the Add Plate window opens.

**Viewing Your User Privileges**

In the User Setup and Preferences window, your privileges are identified by a green check mark in the corresponding checkbox.

- Select the User Setup and Preferences tab.
You can view your assigned privileges at any time, but the check boxes in the display are not enabled unless you are assigned the Add/Manage Users privilege.

Note: Users who are assigned the Add/Manage Users privilege can also create users and update privileges and preferences for any user. Only the superuser can remove the Add/Manage user privilege from a user account.
Table 7 lists the available privileges, which can be assigned in any combination.

### Table 7. User privileges

<table>
<thead>
<tr>
<th>Privilege</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Add/manage users</td>
<td>Add or remove users, set privileges, and change preferences.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Only the superuser can remove this privilege from other users.</td>
</tr>
<tr>
<td>Create new templates</td>
<td>Save a plate or analysis report design as a template.</td>
</tr>
<tr>
<td>View data files (created by other users)</td>
<td>View files created by other users.</td>
</tr>
<tr>
<td>Overwrite existing data file name</td>
<td>Use Save or Save As capability.</td>
</tr>
<tr>
<td></td>
<td>▪ Selecting Save replaces the original file content with any changes made by the user, without changing the file name.</td>
</tr>
<tr>
<td></td>
<td>▪ Selecting Save As allows the user to save either existing or new content with a new file name. Whether or not changes are made, files are automatically saved as files.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Users without this permission can open files and perform analysis but cannot save their changes.</td>
</tr>
<tr>
<td>System settings</td>
<td>View logs, and view and modify the preferred or shared data file and template locations.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> All users can view the file locations.</td>
</tr>
<tr>
<td></td>
<td><strong>Important:</strong> Your system administrator can set preferred locations in System Settings, which override individual user location preferences.</td>
</tr>
<tr>
<td>Maintenance</td>
<td>Perform software updates.</td>
</tr>
<tr>
<td></td>
<td>View the event log.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> All users can view the maintenance log and maintenance reports.</td>
</tr>
<tr>
<td>Data archive</td>
<td>Move raw data from the computer to free up disk space needed for runs on the instrument.</td>
</tr>
</tbody>
</table>
Managing Your Preferences

Use the User Setup and Preferences window to modify your personal user preferences.

Default folder locations appear in your preferences, but you can change the storage locations for your templates and data files. You can also choose to keep all your data files and templates private. The software prompts you to choose either your personal folder or the shared folder each time you save a file or template.

**Important:** If your system administrator enables the Preferred Locations in System Settings, data file folder locations in User Preferences are overridden, and all data files are saved to the preferred file paths. Users can still choose where to store templates.

**To modify your preferences**

1. Tap or click the User Setup and Preferences tab.
2. Change any of the following preferences:
   - Enter a different file path for your data files and templates.
   - Select or clear the check boxes to change your data file and template privacy settings.
     **Note:** The paths and checkboxes are disabled if your administrator has set preferred locations in System Settings.
   - Enter a different system timeout period.
   - Enter a different total of completed plates to show in the Run Status window, up to a maximum of 100.
   - Select the Acquire all Wells By Default checkbox.
     **Note:** If you do not select the checkbox, the default Plate Editor display is all wells excluded from droplet reading and you must manually include the wells for each droplet reading run.
3. Tap or click Save.
4. When the confirmation message appears, tap or click Yes to save the changes, then tap or click OK.

**Signing Out or Changing Users**

While the software is running, one user can sign out and a different user can sign in.

**Note:** Where the screen is locked and the initial user left unsaved changes, an advisory prompt appears. The subsequent user has the option to save or discard the changes before completing the sign-out process.

**To sign out**
1. Tap or click the user name link in the upper-right corner and select Sign Out.
2. Tap or click Yes to confirm.
   - If there are unsaved changes, QX Developer displays a prompt.
     - To discard the changes and proceed, tap or click Yes.
     - To cancel the sign out, tap or click No. Save the changes, and then repeat Steps 1 and 2.

**To change users**
1. Tap or click anywhere in the lockout screen to display the Sign in window.
2. Enter a user name.
   - The domain name appears below the user login fields. If you are on the same domain as the previous user, you can enter your user name only.
     - If the user is on a different domain, enter the domain name followed by a backslash and the user name.
       - `<domain name>\<user name>` *(for example, global\john_smith)*
3. Enter the password and tap or click Sign in.
Instrument Status Bar

An instrument status bar appears above all windows except the analysis displays, which open in a separate module.

When a computer is connected to the QX200 Droplet Reader, the status bar displays the following information:

1. Software name
2. Instrument name and status information
3. Number of runs in progress and runs available
4. Oil and waste levels
   For information on oil and waste level maintenance, see Fluid Maintenance on page 187.
5. Used and free disk space in gigabytes (GB)
   **Note**: When QX Developer detects low disk space, the software displays an advisory prompt to archive data. For information, see Archiving Data on page 199.
6. Name of the current user, along with the current date and time.

If a computer is not connected to the QX200 Droplet Reader, the status bar shows only the software, disk space, and user information.
Functional Windows

This section briefly describes the functional areas in QX Developer. Table 8 explains the primary window accessed from each tab, and includes references to specific chapters or sections in this document where you can find additional information on the functionality.

Table 8. Window Tabs

<table>
<thead>
<tr>
<th>Tab</th>
<th>Name</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Add Plate</td>
<td>Add a plate.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Access the Plate Configuration window, where you can set up your plate for the run. For information on how the functionality is used in this and subsequent windows, see Adding the Plate on page 41,</td>
</tr>
<tr>
<td>Run Status</td>
<td></td>
<td>In the top pane, view the run in process. In the bottom pane, view the completed runs (up to a total of 100). For information on run status, see Running the Experiment on page 54 and Tracking the Run Status on page 55.</td>
</tr>
<tr>
<td>Data Analysis</td>
<td></td>
<td>Access the Data Analysis and Gene Study modules. For information on file analysis, see Data Analysis Module Overview on page 81 and Data Analysis Methodology on page 119.</td>
</tr>
<tr>
<td>Template Setup</td>
<td></td>
<td>Access windows where you can set up plate and report templates. Search for existing template files. Perform lot management for consumables and reagents. For information on creating or editing templates, see Creating or Editing Plate Templates on page 65 and Creating Report Templates on page 154. For information on lot management, see Lot Management for Consumables and Reagents on page 59.</td>
</tr>
</tbody>
</table>
Table 8. Window Tabs, continued

<table>
<thead>
<tr>
<th>Tab</th>
<th>Name</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>System Utilities</td>
<td>Access file storage information, event and maintenance logs, maintenance reports, and tools.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Archive and reprocess data.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>For information, see System Utilities on page 193.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Note: Functionality available to you depends on your assigned user privileges.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Important: Instrument calibration functionality is available only to the Bio-Rad service engineer.</td>
</tr>
<tr>
<td></td>
<td>Users and</td>
<td>View the user privileges you have been assigned, and view and modify your personal preferences.</td>
</tr>
<tr>
<td></td>
<td>Preferences</td>
<td>Create, edit, or remove QX Developer users (if you are assigned the Add/Manage Users privilege).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>For information on your user privileges and preferences, see Getting Started in QX Developer on page 31. For information on setting up users in QX Developer, see Managing Users on page 181.</td>
</tr>
<tr>
<td></td>
<td>Help</td>
<td>Access software version information, the End User License Agreement, the Bio-Rad website, open source software license information, and this document in PDF format.</td>
</tr>
</tbody>
</table>

Compatible File Types

This section describes the file types you can open in QX Developer.

The following applies if your organization is using more than one edition of the software:

- Analysis files created in Regulatory Edition open only in Regulatory Edition.
- Analysis files created in QX Developer open only in QX Developer.
### Table 9. Compatible software file types

<table>
<thead>
<tr>
<th>File Type</th>
<th>Extension</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plate</td>
<td>.ddpitt</td>
<td>Plate template file containing setup details to perform experiments; this file type opens in the Plate Editor.</td>
</tr>
<tr>
<td>Data</td>
<td>.ddpcrd</td>
<td>File type that opens in QX Developer.</td>
</tr>
</tbody>
</table>
Chapter 6  Running Experiments

After the QX Developer recognizes the plate holder inserted in the QX200 Droplet Reader, you can open the Plate Configuration window where you can identify your plate, configure your plate layout, and start the run. The software enables each button sequentially (Add Plate, Configure Plate, Start Run) when you complete the prerequisite tasks.

Adding the Plate

This section explains how to add the plate in QX Developer. You can insert a plate into the instrument before or after you add the plate in the software. For information on how to properly insert a plate into the QX200 Droplet Reader, see Inserting an Assembled Plate into the Instrument on page 27.

To add a plate in QX Developer

1. Tap or click the Add Plate tab.

If the Runs Available number in the Instrument Status Bar is 1, the Add Plate button is enabled.

**Note:** You cannot add a plate in the software if the QX200 Droplet Reader is currently processing a run.

2. Tap or click next to Add Plate.

When QX Developer recognizes the plate holder in the compartment, the software enables the Configure Plate button.

**Note:** If QX Developer does not recognize the plate holder, the software displays an advisory prompt.

3. Tap or click Configure Plate.

The Plate Configuration window opens.
Plate Configuration Window

After you tap or click Configure Plate, the Plate Configuration window opens to the Plate Information display.

All users can start runs on selected plates using the initial configuration, an altered configuration, or just the plate and data file names and the supermix.

When configuring plates

- Users without the Create New Template user privilege can customize a plate design before starting a run but cannot save the file.
- Users with the Create New Template user privilege can save a new or modified plate design as a template before starting the run.

Plate Information Tab

The Plate Information tab is selected by default when the Plate Configuration window opens, and provides functionality allowing you to quickly set up a plate for a run.

You can

- Select, modify (optional), and run an existing template
- Create and run a new plate design
- With the applicable user privilege, save existing or new plate designs as templates
- Identify a plate name, a supermix, and a data file name (optional) for a run

**Note:** At a minimum, you must enter a plate name and select a supermix to enable the Start Run button. The data file name field is automatically populated with the plate name, but you can change it. You can identify the remainder of your experiment parameters either in the Plate Editor before you start the run, or in the plate editor layout in the Analysis module after the run has concluded.

- Select the well acquisition method the instrument should use
Configuring the Plate

To select an existing plate template

1. Under Plate Template, tap or click the dropdown arrow under Choose Existing and select a template file. The software automatically populates the fields under Create New Plate.

2. (Optional) Under Create New Plate, change the plate or data file name.

3. (Optional) Under Acquire Wells By, select the alternative well acquisition method. The default acquisition is by column.

4. Tap or click the Well Selection tab and ensure all empty wells are excluded from droplet reading. For information, see Including or Excluding Wells in the Plate Layout on page 45.

5. Select the Well Information tab, and then select the wells to be processed in the run.

6. (Optional) Change the configuration for one or more wells. For information, see Defining or Editing Well Information on page 47.

7. Click Apply.

The Start Run button is enabled.
To create a new template

1. Under Plate Template, tap or click Create New to open the corresponding editor window in template Edit mode.

   **Important:** You must be assigned the Create New Template user privilege, which allows you to save the template. Otherwise, you must use an existing template or use the Create New Plate option and configure your plate from the Well Selection and Well Information tabs.

   The Plate Editor opens in Edit mode.

2. Click the Exclude button and include the wells to be processed. For information, see Including or Excluding Wells in the Plate Layout on page 45.

   By default, all wells in the Plate Editor are automatically excluded, and you need to enable the wells to be run. If you typically run full plates, you can change the setting to Acquire all wells by default in your user preferences. For information, see Managing Your Preferences on page 35.

   **Important:** All empty wells must be excluded.

3. Configure experiment parameters for each well to be processed. For information see Creating or Editing Plate Templates on page 65.

4. (Optional) Under Acquire Wells By, select the alternative well acquisition method. The default acquisition is by column, but you can direct the instrument to acquire wells by row.

5. Click Apply.

6. Click Save and select one of the following:

   - Shared Templates. The template is saved in the folder specified in the System Settings.
   - My Templates (saves the file to the path in your user preferences) or System Templates (saves the file to the preferred location specified by your system administrator).
   - If your system administrator has specified preferred locations for all users, your template must be saved to that location.

   The Start Run button is enabled.

To run a plate without defining parameters

1. Enter a plate name under Create New Plate.

2. Select a supermix.

3. Enter a file name for the data file that QX Developer generates at the end of the run.

   The Start Run button is enabled.
Well Selection Tab

When you select the Well Selection tab, a blank Plate Editor grid appears. The QX200 Droplet Reader requires empty wells to be excluded from droplet reading; therefore, the QX Developer Plate Editor layout shows all wells as excluded by default, and you must select and manually include the wells to be run.

Tip: If you typically run full plates, you can change the setting in User Preferences to acquire all wells by default. For information, see Managing Your Preferences on page 35.

Including or Excluding Wells in the Plate Layout

To identify wells to include in the experiment run

1. Select the Well Selection tab.
2. In the Plate Editor layout, select the wells containing sample.
3. Tap or click Include Selected Wells.
On the right, QX Developer specifies the number of wells that will be included in the droplet reading run.

**To identify wells to exclude in the experiment run**

4. Select the Well Selection tab.
5. In the Plate Editor layout, select all empty wells.

6. Tap or click Exclude Selected Wells.

On the right, QX Developer specifies the number of wells that will be included in the droplet reading run.
**Well Information Tab**

When you select the Well Information tab, the Plate Editor opens.

**Note:** The window is identical to the Edit window in Template Setup.

You can configure your experiment parameters for selected wells before or after the run. However, you must select a supermix, and identify the plate and file name before the Start Run button is enabled.

**Defining or Editing Well Information**

You can assign different experiment parameters to each well or groups of wells. You must tap or click Apply for the software to recognize your new or edited information.

You can apply your entries at any time.

**Note:** If you tap or click the Save button before the plate layout is complete, the file closes and you must reopen it from the saved template location.
To set up your wells in the plate

1. If you are accessing the Plate Editor from
   - the Add Plate tab, tap or click Configure Plate and select the Well Information tab to display the Plate Editor.
   - the Template Setup tab, the Plate Editor opens with the Edit button selected by default.
   If you are creating a new template, the grid is blank. If you are editing an existing template, information appears for the configured wells.

2. Select a well or group of wells.

3. Select an experiment type. For more information, see Experiment Types on page 73.

4. In the Sample Description fields, enter up to four words or phrases that describe the sample. For more information, see Sample Descriptions on page 74.

5. Select a sample type. For more information, see Sample Types on page 75.

6. Select a supermix. For more information, see Supermixes on page 75.

7. Select an Assay Type. Available assay types vary, depending on the selected Experiment Type. For information, see the fluorophore options in Assay Types and Fluorophores on page 76.

8. (Optional) Under Target Info, change the fluorophore assigned to a channel.
   
   The fluorophore information populates automatically for each target, with the maximum number of rows allowed for the assay method, but you can modify the default fluorophore assignments, if applicable.
   
   a. Tap or click the field dropdown arrow and select an alternative.
   b. To delete a row, tap or click the minus (-) icon next to the target.
   c. To re-add a row, tap or click the plus (+) icon.

9. (Optional) To clear one or more wells at any time, select the wells and tap or click Clear Selected Wells.
10. Tap or click Apply.

**Tip:** After you have applied your entries and selections, you can pause the cursor on a well to see the well information.

11. If you are finished with your plate configuration, tap or click Save.
Starting the Run

When all required elements to set up a plate for a run in QX Developer are satisfied, the software enables the Start Run button.

**Important:** Ensure there is sufficient QX200 Droplet Reader Oil available, and the waste bottle is less than 70% full.

![Screenshot of QX Developer software interface showing the Start Run button.]

- Tap Start Run.

  To reauthenticate your user credentials, you are prompted to sign in again.

  Enter your user name and password, and then tap or click Sign in.

  The software performs a mandatory prerequisites check to ensure that the instrument is ready.

  One of the following occurs:

  - If the instrument passes the check, the run starts normally. Continue to [Running the Experiment on page 54](#).
  - If the check finds anything amiss on the instrument, the software displays a dialog box in which the issues are identified. You must resolve each issue before continuing. For information, see [Run Prerequisites Checklist](#).

![Screenshot of a required consumables check for run dialog box.]

The instrument cannot start the run until all specified issues are resolved.
Run Prerequisites Checklist

If the prerequisites check on the instrument status finds one or more problems with instrument readiness, QX Developer. identifies them with an ❌ in the Check for Run dialog box. You must resolve all issues before the instrument can start the run.

**Note:** If you encounter a problem with the instrument or software that is not described here, see Troubleshooting on page 205.

For information on each issue that can prevent the instrument from starting the run, see Table 10.
Table 10. Pre-Run Error Indicators

<table>
<thead>
<tr>
<th>Status Check Item</th>
<th>Issue</th>
<th>Suggested Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Instrument Connection</td>
<td>USB cable connection not recognized</td>
<td>Disconnect, and then reconnect, the USB cable between the instrument and the computer.</td>
</tr>
<tr>
<td></td>
<td>Power connection not recognized</td>
<td>Ensure the power outlet is working. Disengage the power cord from the outlet, and then reinset it into the outlet.</td>
</tr>
<tr>
<td>Fluid Levels Stability</td>
<td>The oil or waste is moving within the bottle so the level cannot be accurately determined.</td>
<td>Wait until the liquid is settled, then try again. Ensure the instrument rests on a solid, stable surface.</td>
</tr>
<tr>
<td>Sufficient Oil</td>
<td>Oil level is below the acceptable level for a run.</td>
<td>Remove the oil bottle and replace it with a full one.</td>
</tr>
<tr>
<td>Sufficiently Low Waste</td>
<td>Waste level is above the acceptable level for a run.</td>
<td>1. Empty any remaining oil from the bottle and then place a Waste label (provided by Bio-Rad) over the Oil label.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Insert the bottle into the Waste slot.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>For information on changing containers, see Instrument Maintenance on page 187.</td>
</tr>
<tr>
<td>Bottle Door Closed</td>
<td>Bottle door is not closed properly.</td>
<td>Open the bottle door and ensure there are no blockages, and then close the door again.</td>
</tr>
<tr>
<td>Reader Door Closed</td>
<td>Reader door is not closed properly.</td>
<td>Open the reader door and ensure there are no blockages, and then close the door again.</td>
</tr>
<tr>
<td>Reader Lid Closed</td>
<td>Reader lid is not closed properly.</td>
<td>Open the reader lid and ensure there are no blockages, and then close the lid again.</td>
</tr>
</tbody>
</table>
### Table 10. Pre-Run Error Indicators, continued

<table>
<thead>
<tr>
<th>Status Check Item</th>
<th>Issue</th>
<th>Suggested Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plate Loaded</td>
<td>Plate is not loaded properly.</td>
<td>1. Open the reader door and remove the plate.&lt;br&gt;2. Ensure the tabs of the plate cover are securely clamped.&lt;br&gt;3. Insert the plate again and ensure it lays flat.&lt;br&gt;For information on loading plates in the QX200 Droplet Reader, see Inserting an Assembled Plate into the Instrument on page 27</td>
</tr>
<tr>
<td>Sufficient Disk Space</td>
<td>There is not enough storage space available for the run data.</td>
<td>Archive older data files to clear space.&lt;br&gt;For information on archiving data files, see Archiving Data on page 199.</td>
</tr>
</tbody>
</table>

After you resolve the issues, continue to Running the Experiment.
Running the Experiment

During and after the run, QX Developer displays run information in the Run Status window. For information, see Tracking the Run Status.

- During the run, you can perform real-time analysis on the droplets in each well after the well has been read. For information, see Using Live Analysis on page 56.
- You can cancel the run at any time. For information, see Canceling a Run on page 58.
- When the run concludes, QX Developer moves the run to the Completed list in the Run Status window, and saves the analysis data file.

Data files are always saved to one of the following locations:

- Your personal file location specified in your user preferences
- The preferred location in System Settings

Important: if your system administrator has designated preferred locations for all users, these file paths override the file paths indicated in your preferences.

For information on file storage locations, see System Utilities on page 193. For information on opening and analyzing your data file, see Data Analysis Module Overview on page 81.
Tracking the Run Status

You can track each run from the Run Status window.

The top pane shows the run in progress. The run remains in the top pane for the duration of the processing.

Icons displayed in the Run Status column indicate each process point, as shown in Table 11.

**Table 11. Run status indicators**

<table>
<thead>
<tr>
<th>Status Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ready to begin droplet reading</td>
</tr>
<tr>
<td></td>
<td>Droplet reading</td>
</tr>
<tr>
<td></td>
<td>Run completed, plate in Outbox</td>
</tr>
<tr>
<td></td>
<td>Run completed, plate removed from Outbox</td>
</tr>
<tr>
<td></td>
<td>Run stopped or canceled by the system</td>
</tr>
</tbody>
</table>

When the run is finished, the software moves the run to the bottom pane, where up to 100 completed runs are visible.

**Tip:** You set the number of completed runs to display in your user preferences. See Managing Your Preferences on page 35.
The content of each column is explained in Table 12.

### Table 12. Run Information Columns

<table>
<thead>
<tr>
<th>Column Name</th>
<th>Location</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plate name</td>
<td>Top and bottom panes</td>
<td>Name entered under Create New Plate</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Note:</strong> If you selected an existing template, the template name appears.</td>
</tr>
<tr>
<td>Run status</td>
<td>Top and bottom panes</td>
<td>Icons for each process step, as explained in Table 11 on page 55</td>
</tr>
<tr>
<td>Time remaining</td>
<td>Top pane only</td>
<td>Hours and minutes until the run will be finished</td>
</tr>
<tr>
<td>Time elapsed</td>
<td>Top pane only</td>
<td>Time, in hours and minutes the run has been in progress</td>
</tr>
<tr>
<td>Total time remaining</td>
<td>Below the top pane</td>
<td>Total hours and minutes remaining for the run</td>
</tr>
<tr>
<td>Run completed</td>
<td>Bottom pane only</td>
<td>Date and time stamp showing when the instrument completed the run</td>
</tr>
<tr>
<td>File name</td>
<td>Bottom pane only</td>
<td>Plate name and system-generated identifier</td>
</tr>
</tbody>
</table>

After run completion, the software saves the analysis data file in either your personal file location or to a preferred location specified by your system administrator.

**Important:** If your system administrator has specified preferred locations for all users in System Settings, files are saved there, rather than to the path specified in user preferences. For information, see Preferred Location on page 194.

If your system is connected to a network, and shared folders are identified in the global preferences, the data file is also copied to a shared folder.

### Using Live Analysis

When the QX200 Droplet Reader initiates droplet reading, QX Developer provides an option for real-time analysis of the wells that have been read, while the process as a whole is still active.

QX Developer initially shows each well in the Plate Editor as disabled. As the software confirms that the instrument has acquired a well and read the droplets, the software enables the well for live analysis in read-only mode. The process continues until all wells are enabled and the process is finished.
To use live analysis

1. Tap the droplet reading icon (●) for the run in progress.
   The Droplet Reading dialog box opens.

2. Tap Open Live Analysis.
3. Select an analysis window.
4. In the well selector, select one or more wells that are enabled.

   **Important:** QX Developer displays read-only data in the selected analysis window. You can view the data but to make changes, you must wait for the run to complete, and then view the data in the Analysis module.
Canceling a Run

You can cancel a run at any time during the droplet reading process.

1. In the Run Status window, tap or click the droplet reading icon (在校) for the run to be canceled.

   The Droplet Reading dialog box opens.

   ![Droplet Reading dialog box]

2. Click Abort Plate Run.

3. When the confirmation message appears, tap or click Yes.

   The instrument finishes reading the current well, stops the process, and reverts to its pre-run state.

   The icon appears in the Run Status column.

   QX Developer then creates the data file for the wells that were processed, and the run end status in the Event Log is specified as “run aborted by user.”

   **Note:** If you tap or click No, or if you do not respond to the confirmation message within 30 seconds, the process continues without interruption.
Chapter 7 Lot Management for Consumables and Reagents

You can maintain an inventory and track the use of your consumables and reagents in the QX Developer software.

For consumables, you can manage lots of the following items:

- ddPCR Dx AutoDG consumables packs
- Cartridges
- Plates and seals
- Pipet tips

For reagents, you can manage lots of the following items:

- Supermix
- Droplet generator oil
- Droplet reader oil
- Buffer controls
- Assays

Lots that have been fully used are not displayed.
Managing Consumable Lots

To enter a consumables lot

1. Select the Template Setup icon.

2. Select Lot Management, and then select the Consumables tab.

   From the Consumables window, the following tabs appear above the lot information grid:
   - ddPCR Dx AutoDG Consumables Pack (appears by default)
   - Cartridges
   - Plates and Seals
   - Pipet Tips

3. Select a tab.

4. (Optional) To display expired or used lots, select the checkbox at the bottom-left side of the window.

5. Click in each field and enter the corresponding information as follows:
   - Lot Number
Managing Reagent Lots

- Catalog Number
- Lot Expiry Date

6. If the consumable was used in a run, select the Used checkbox.
7. Repeat until all lots are added.
8. Tap or click Save.

Managing Reagent Lots

To enter a reagent lot

1. Select the Template Setup icon.

2. Select Lot Management, and then select the Reagents tab.

   From the Reagents window, the following tabs appear above the lot information grid:
   - Supermix (appears by default)
   - Droplet Generator Oil
   - Droplet Reader Oil
Chapter 7 Lot Management for Consumables and Reagents

- Buffer Controls
- Assays

3. Select a tab.

4. (Optional) To display expired or used lots, select the checkbox at the bottom-left side of the window.

5. Click in each field and enter the corresponding information as follows:
   - Supermix (in Supermix grid only)
   - Lot Number
   - Catalog Number
   - Lot Expiry Date

6. If the consumable was used in a run, select the Used checkbox.

7. Repeat until all lots are added.

8. Tap or click Save.

**Showing Lots in Run Information Output and Reports**

After you have set up your lots, you can ensure that lots appear in the Run Information screen of your analysis files.

**To show the lots in Run Information**

1. Click Add Plate, then click Configure Plate.
2. Select the Lot Selector tab.
3. For consumables and reagents
   a. Select each applicable lot for the run
   b. After each selection, click Select Lots.
4. Continue setting up the plate until the Start Run button is enabled.
5. Click Start Run.
6. When the run is completed, open the analysis file.
7. Click the Run Information tab to see the used lots.
To show the lots in a report

► Ensure the Lot Information check box is selected in the report elements before running the report.
Chapter 8 Creating or Editing Plate Templates

If you are assigned the Create New Templates user privilege, you can save a new or reconfigured plate as a template for reuse. Templates that are available to you appear in the default view of the Template Setup tab, as follows:

- Templates you created and saved, which are automatically saved to one of the following paths:
  - The path specified in your user preferences; these appear under My Templates
  - The path specified under Preferred Locations in System Settings, if your system administrator has specified the path for all users, these appear under System Templates

  **Important:** If your administrator has specified the Preferred Location for all users, the path overrides the path specified in user preferences. Preferred locations can be enabled only by users with the System Settings user privilege.

- Templates created and saved by any user and designated as shared; these appear under Shared Templates.

From the Template Setup or Add Plate window, you can create or modify plate templates. From the Template Setup window, you can also modify reports.
Plate Editor Window

Use the Template Setup Plate Editor to

- Create new or edit existing plate template files
- Save new or modified templates for reuse (with Create New Templates permission only)
- Define or change experiment type, sample type, supermix, and assay type for each well
- Set reference targets and control samples
- Exclude wells from droplet reading
- Add applicable plate or well notes

The Plate Layout view is the default template view.

**LEGEND**

1. You can access the Edit and Exclude buttons from the top of the screen.

2. The right pane contains the interface for defining your experiment parameters.
**Legend, continued**

3. The left pane displays the plate grid and configuration information in each well.

4. The Well Data button toggles to the Well Data table, where you can see your plate setup in a tabular format, and then back to the Plate Editor. When the Well Data table is displayed, the button name changes to Plate Layout.

![Plate Layout](image)

**Plate Editor Tools**

Use the buttons described in Table 13 to define your experiment in the Plate Template Editor.

<table>
<thead>
<tr>
<th>Button</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Edit</td>
<td>Allows you to</td>
</tr>
<tr>
<td></td>
<td>- Select wells or clear well selections</td>
</tr>
<tr>
<td></td>
<td>- Set the experiment type, sample name and type, supermix, and assay type</td>
</tr>
<tr>
<td></td>
<td>- Set the target name, type, and fluorophore for each channel</td>
</tr>
<tr>
<td></td>
<td>- Add well or plate notes</td>
</tr>
<tr>
<td></td>
<td><strong>Note</strong>: This button is enabled by default when you open the window.</td>
</tr>
<tr>
<td>Exclude</td>
<td>Allows you to exclude wells from droplet reading</td>
</tr>
<tr>
<td>Cancel</td>
<td>Allows you to cancel and return to the Template Setup window</td>
</tr>
<tr>
<td>Save</td>
<td>Opens a dialog box where you can enter a file name and specify a storage location</td>
</tr>
<tr>
<td>Clear Selected Wells</td>
<td>Allows you to delete information from selected wells</td>
</tr>
<tr>
<td>Undo</td>
<td>Allows you to reverse your last action</td>
</tr>
<tr>
<td></td>
<td><strong>Note</strong>: This button is enabled only after you have applied at least one experiment parameter (for example, the selection for Experiment Type).</td>
</tr>
<tr>
<td>Well Data or Plate Layout</td>
<td>Toggles button to access the Well Data table and return to Plate Layout view</td>
</tr>
<tr>
<td>Apply</td>
<td>Applies your entries and choices for selected wells</td>
</tr>
</tbody>
</table>

**Table 13. Plate Editor Buttons**
Opening a New Plate Template

You can open a blank plate template from either the Template Setup window or the Add Plate window.

- Use Template Setup if you are creating one or more templates to store for future use.
- Use Add Plate if you are creating a template to correspond to a plate you are about to run.

To open a new plate template from the Template Setup window

1. Tap the Template Setup tab.

2. Tap Plate Setup.

3. Tap Add New Plate.

4. Tap Edit Plate Layout.
   
   The Plate Editor opens, displaying a blank plate template.

5. To set up your plate, continue to Defining or Editing Well Information on page 47.

To open a new plate template from the Add Plate window

1. Tap the Add Plate tab.

2. Tap or click the + icon next to Add Plate.

3. When the software enables the Configure Plate button, tap or click Configure Plate.
   
   The Plate Configuration window opens to the Plate Information tab by default.

4. Under Plate Template, tap or click Create New.
   
   The template Plate Editor opens, displaying a blank plate layout.

5. To set up your plate, continue to Defining or Editing Well Information on page 47
Opening an Existing Plate Template

You can open an existing plate template from either the Template Setup window or the Add Plate window.

- Use Template Setup if you are editing one or more templates to store for future use.
- Use Add Plate if you are editing a template to correspond to a plate you are about to run.

To open an existing plate file from the Template Setup window

1. Tap or click the Template Setup tab.

   ![Template Setup Tab]

   The Plate Setup window appears by default and a list of available plate templates appears.

2. Select a template and tap or click Edit Plate Layout.

   **Tip:** You can also double-click the template file to open it.

   The Plate Editor opens, displaying the plate template as it is currently configured.

   To make changes to the plate, continue to Defining or Editing Well Information on page 47

To open an existing plate template from the Add Plate window

1. Tap the Add Plate tab.

   ![Add Plate Tab]

2. Tap or click the + icon next to Add Plate.

3. When the software enables the Configure Plate button, tap or click Configure Plate.

   The Plate Configuration window opens to the Plate Information tab by default.

4. Under Plate Template, tap or click the dropdown arrow and select a template from the list.

   The template Plate Editor opens, displaying the configured layout.

   You can run the plate without changing the template, or you can modify the configuration. To make changes to the plate, continue to Defining or Editing Well Information on page 47
Template Setup — Setting Up Your Wells

When you open the Plate Editor from the Template Setup tab, the Edit button is selected by default. You can assign different experiment parameters to each well or groups of wells.

For each well, or group of wells, you must

- Choose an experiment type
- Select a sample type
- Select a supermix
- Select an assay method

Optionally, you can

- Enter up to four sample descriptors for each well
- Modify fluorophore assignments
- Add notes for each well
- Add notes for the plate

If you are creating a new template, the grid is blank. If you are editing an existing template, previously saved information appears when you select one or more configured wells.
To define well parameters

1. Select a well or group of wells.

2. From the Experiment Type dropdown list, select a new or different experiment type. For more information, see Experiment Types on page 73.

3. In the Sample Description fields, enter or change up to four words or phrases that describe the sample. For more information, see Sample Descriptions on page 74.

4. From the Sample Type dropdown list, select a new or different sample type. For more information, see Sample Types on page 75.

5. From the Supermix dropdown list, select or change the supermix. For more information, see Supermixes on page 75.

6. The Assay Type field populates automatically based on your experiment selection. To change it, select a different assay type from the dropdown list. For information, see the fluorophore options in Assay Types and Fluorophores on page 76.

   Note: Available assay types vary, depending on the selected experiment type.

7. (Optional) Under Target Info, change the fluorophore assigned to a channel.

   The fluorophore information populates automatically for each target, with the maximum number of rows allowed for the assay method, but you can modify the default fluorophore assignments, if applicable.

   a. Tap or click the field dropdown arrow and select an alternative.

   b. To delete a row, tap or click the minus (-) icon next to the target.

   c. To re-add a row, tap or click the plus (+) icon.

8. (Optional) To clear one or more wells at any time, select the wells and tap or click Clear Selected Wells.

9. Tap or click Apply.

   Note: You must tap or click Apply for the software to recognize your new or edited information, and before you can save the template. Once applied, you can pause on a well to see the corresponding information in a tool tip.
10. Ensure empty wells are excluded from droplet reading. For information, see Template Setup – Including or Excluding Wells on page 78.

11. If you are finished with your plate configuration, tap or click Save in the upper-right corner, and then in the Save dialog box, enter a new name for the template and tap or click Save again.

   **Note:** You must save the template with a new name. You cannot overwrite existing templates. If you tap or click the Save button before the plate layout is complete, the file closes and you must reopen it from the saved template location.
Experiment Types

QX Developer offers seven experiment types from which you can choose, each enabling different assay options and analysis tools. Table 14 briefly explains each experiment type.

Tip: You can assign multiple experiment types within the plate, but you can assign only one experiment type to each well.

Table 14. Experiment types

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direct quantification (DQ)</td>
<td>DQ uses absolute quantification to determine the concentration (copies/µl) of target DNA copies in a sample. DQ measures the number of positive and negative droplets for each target in a well based on their fluorescence amplitude, and uses a Poisson algorithm to calculate the starting concentration of each target DNA molecule. <strong>Note:</strong> The ABS experiment type created in QuantaSoft Software versions 1.4 through 1.7 is automatically mapped into the software as a DQ experiment type.</td>
</tr>
<tr>
<td>Copy number variation (CNV)</td>
<td>CNV determines concentration and calculates the copy number of an unknown (CNV) target relative to a known reference or references within the same well.</td>
</tr>
<tr>
<td>Mutation detection (MUT)</td>
<td>MUT determines concentration and calculates the fractional abundance of an unknown mutant present at a low frequency in a wild-type background.</td>
</tr>
<tr>
<td>Rare event detection (RED)</td>
<td>RED determines the concentration of a known mutant or a rare target species relative to a given reference species within a large pool of background DNA.</td>
</tr>
<tr>
<td>Drop-off (DOF)</td>
<td>DOF determines absolute quantification of targets for assays designed to detect non-wild type sequences, such as indels and genome edits; DOF is designed to support an assay strategy where one probe counts all alleles and one “drop-off” probe sits on top of a predicted cut site.</td>
</tr>
<tr>
<td>Gene expression (GEX)</td>
<td>GEX determines concentration (as with DQ) and calculates relative expression levels of an unknown target relative to a known reference or references within the same well.</td>
</tr>
<tr>
<td>Residual DNA quantification (RDQ)</td>
<td>RDQ provides a precise method for residual host cell DNA quantification. <strong>Note:</strong> Users can enter a conversion factor (1/C-value) for a given genome (human, mouse, etc.) to calculate the mass inputted into a well based on the number of copies measured, up to three decimals. The software displays the result in pg/µl of the ddPCR well in the Molecular Weight column of the Data Table. Starting sample concentration can then be back-calculated based on dilution factor.</td>
</tr>
</tbody>
</table>
Sample Descriptions

QX Developer allows you to enter descriptive words or phrases for your sample in up to four fields per well. Descriptors can include information such as research type, dilution factors, and so forth.

Tip: Only the information from the first field appears in the Plate Editor layout. However, you can see entries in the second, third, and fourth fields in the Well Data table, or you can pause on a well to display a tool tip, as shown in the following graphic.
Sample Types

QX Developer offers four sample types:

- Unknown
- NTC (no template control)
- Positive control, where one or more targets are expected
- Negative control, where no response is expected

Tip: You can assign multiple sample types within the plate, but you can assign only one sample type to each well.

Supermixes

Important: Use of unapproved supermixes can harm the instrument and void the warranty.

QX Developer offers five PCR supermixes. Each supermix is optimized to deliver maximum PCR efficiency and sensitivity for the amplification and detection of DNA and RNA targets.

- **ddPCR Multiplex Supermix** — For use in nucleic acid sample preparations to amplify and detect DNA targets in probe-based experiments involving multiple targets
- **ddPCR Supermix for Probes (No dUTP)** — For use in nucleic acid sample preparations, with sensitivity for the amplification and detection of DNA targets using hydrolysis probe-based assays
- **ddPCR Supermix for Probes** — For use in sample preparation with uracil N-glucosylase (UNG) decontamination protocols to prevent the reamplification of carryover PCR products between experiments
- **ddPCR Supermix for Residual DNA Quantification** — For use in residual DNA detection
- **One-Step RT ddPCR Advanced Kit for Probes** — For use in absolute quantification of target RNA molecules
Assay Types and Fluorophores

QX Developer offers a total of five assay methods, which are explained in Table 15.

Table 15. Assay types

<table>
<thead>
<tr>
<th>Assay type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single target per channel</td>
<td>This assay type is available for the DQ, CNV, MUT, and GEX experiment types.</td>
</tr>
<tr>
<td>Amplitude multiplex</td>
<td>Method to increase multiplexing up to four targets per well, with one or two targets detected per channel. The maximum of eight rows are displayed. You can delete up to seven rows.</td>
</tr>
<tr>
<td>Probe mix triplex</td>
<td>Triplex mode that allows exactly three targets per well, with one target detected in each of two channels and a third target detected across both channels.</td>
</tr>
</tbody>
</table>

**Note:**

Advanced classification | Method to increase multiplexing rate up to targets per well in any combination of method targets detected per channel. This method requires manual classification of droplets. This assay type is available only for the DQ experiment type. |
| Basic drop-off          | This mode allows two groups of two targets each. Group 1 is shown by default, with one target in FAM and HEX/VIC and one target in FAM or HEX/VIC. This assay type is available only for the DOF experiment type. |
The experiment type you select determines the available assay methods. Table 16 displays the sample type associated with each assay method.

**Table 16. Assay types by experiment type**

<table>
<thead>
<tr>
<th>Assay type</th>
<th>DQ</th>
<th>CNV</th>
<th>MUT</th>
<th>DOF</th>
<th>GEX</th>
<th>RDQ</th>
<th>RED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single target per channel</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Amplitude multiplex</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Probe mix triplex</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Advanced classification</td>
<td>✓</td>
<td></td>
<td></td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Basic drop-off</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

QX Developer automatically populates the fluorophore in each channel for each target but you can change the default selections.

Use Table 17 to select from the fluorophore options for each assay method listed.

**Table 17. Fluorophore options**

<table>
<thead>
<tr>
<th>Assay method</th>
<th>Signal Ch1</th>
<th>Signal Ch2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single target per channel, for 1 to 2 targets</td>
<td>FAM</td>
<td>HEX</td>
</tr>
<tr>
<td></td>
<td>EvaGreen®</td>
<td>VIC</td>
</tr>
<tr>
<td></td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Amplitude multiplex, for 1 to 4 targets</td>
<td>FAM Hi</td>
<td>HEX Hi</td>
</tr>
<tr>
<td></td>
<td>FAM Lo</td>
<td>HEX Lo</td>
</tr>
<tr>
<td></td>
<td>EvaGreen® Hi</td>
<td>VIC Hi</td>
</tr>
<tr>
<td></td>
<td>EvaGreen® Lo</td>
<td>VIC Lo</td>
</tr>
<tr>
<td></td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Probe mix triplex, for 3 targets</td>
<td>FAM</td>
<td>HEX</td>
</tr>
<tr>
<td></td>
<td>EvaGreen®</td>
<td>VIC</td>
</tr>
<tr>
<td></td>
<td>None</td>
<td>None</td>
</tr>
</tbody>
</table>
Template Setup – Including or Excluding Wells

Optionally, you can include or exclude wells from droplet reading.

**To identify wells to include in the experiment run**

1. Select the Well Selection tab.
2. In the Plate Editor layout, select the wells containing sample.
3. Tap or click Include Selected Wells.

On the right, QX Developer specifies the number of wells that will be included in the droplet reading run.

**To identify wells to exclude in the experiment run**

4. Select the Well Selection tab.
5. In the Plate Editor layout, select all empty wells.
6. Tap or click Exclude Selected Wells.

On the right, QX Developer specifies the number of wells that will be included in the droplet reading run.
Chapter 8 Creating or Editing Plate Templates
Chapter 9 Data Analysis Module Overview

As the instrument reads droplets in each well, the QX200 Droplet Reader collects raw data. At the end of each droplet reading run, QX Developer uses the raw data to produce a data file, which contains analysis information based on the number of positive and negative droplets identified for each target in each sample. The fraction of positive droplets is fit to a Poisson algorithm to determine the starting concentration of the target DNA molecule in units of copies/µl input.

When you open the data file in the Analysis module, you can view the data with the automatic analysis thresholds and calculations, or you can choose different options to recalculate, replot, or recolor the results.

When you select the Data Analysis tab, you can access two modules:

- ddPCR Data Analysis
- Gene Study Analysis

The Data Analysis module functionality allows you to

- View your droplet data in 1D and 2D plots
- Analyze multiple targets within a well using amplitude multiplexing and probe mixing strategies
- Detect genomic editing events and non-wild type events with a “drop-off assay” analysis option
- Visualize and export data with improved flexibility
- Generate reports for different analysis types
Chapter 9 Data Analysis Module Overview

The Gene Study module allows you to

- Collect data by adding one or more data files
- View gene expression calculations across multiple experiments

**Note:** The maximum number of samples you can analyze in a gene study is limited by the size of the computer’s RAM and virtual memory.

More detailed information, including specific information and uses for each analysis window, as well as information on data calculations, is contained in.

## Opening a Datafile for Analysis

You can open the following file types in the Data Analysis module:

- .qlp
  
  If you open this file type, the Plate Editor appears by default. When you save this file type in QX Developer, the software automatically saves the file as a .ddpcr.ddpcri.ddpcrd file. This creates a new file and does not overwrite the original.

- .ddpcrd

Analysis data files are typically stored in one of the following defined storage locations:

- Personal folder for the individual user on the connected computer or the system folder specified for Preferred locations in the System Settings window
  
  **Note:** Your personal folder location is displayed in your user preferences. You can open the System Settings window to see the preferred or shared storage locations.

  **Important:** If your system administrator has specified preferred location file paths for all users, the file paths specified in user preferences are overridden.

- Shared folder in a different location, such as a network path
  
  **Note:** The software also saves data files to a shared folder if it is accessible.

To open data files, you must be on a computer on which QX Developer is installed and have access to the storage locations.

**To open a data file for analysis**

1. Open QX Developer and log into the application.
2. Select the Data Analysis tab.
Available data files are displayed under the following headings.

- **Recent Datafiles**
  Displays a list of files you have recently opened.

- **System Datafiles or My Datafiles**
  Only one of the following locations is available:
  - If your system administrator has enabled Preferred Locations for all users, all data files are stored in that folder, and files appear under System Datafiles.
  - If preferred locations are not enabled for all users, your data files are saved to your personal folder and appear under My Datafiles.

- **Shared Datafiles**
  Displays a list of data files saved to the shared location, if one has been specified in the System Settings window.

  **Important:** Data files are always saved to either My Datafiles or System Datafiles. Data files are also saved to the Shared location (such as a network directory) if it is accessible when the software saves the data file.

3. Tap or click a file to open it in the Analysis module.
   
   Tabs for each analysis and data window appear on the left.

   **Tip:** From the computer attached to the QX200 Droplet Reader, you can also open the data file from the Run Status window. In the completed files section, tap or click the completed run icon (✔️) for the file, and then tap or click Launch Analysis.
Saving a Datafile

You can save data files with or without modifications, with the same or different file names, and in the same or different locations.

**Note:** To save a file with the same file name (Save button), you must be assigned the Overwrite Existing Datafile name user privilege.

**To save a data file**

1. Do one of the following:
   - Tap Save to save the file under the same file name and in the same location.
   - Tap Save as to save the file under a different file name or in a different location.
     - If you are saving the file under the same name and in the same location, QX Developer saves the file immediately.
     - If you are saving the file under a different file name, or in a different location, continue to Step 3.

   To reauthenticate your user credentials, you are prompted to enter a Save Reason, and then sign in again.

2. Enter the Save Reason, and then tap or click Save As.

3. Enter your user name and password, and then tap or click Sign in.
   - If you are saving the file under the same name and in the same location, QX Developer saves the file immediately.
   - If you are saving the file under a different file name, or in a different location, continue to Step 3.

4. Navigate to the new location and/or enter a new file name.

5. Tap Save.
Analysis Dashboard

When you open a data file, the Analysis module opens to a Dashboard window containing default analysis and data windows in a summary view. You can customize the view to increase or decrease the number of windows that appear in the Dashboard, and you can also change the selection of windows that appear.

For information on the available options in the Dashboard view, see Dashboard Options on page 87.

Note: If you open a .qlp file, the Plate Editor opens as the default view and you must select the Dashboard tab to open it.

From the tabs on the left, you can access plate views, amplitude charts, statistical probability distribution charts, droplet counts, and a complete data table. You can also view data specific to the run in the Run Information window, and run reports on any of the analysis charts from the Reports window.
Modifying the Dashboard Display

You can modify the column number and row number to display any or all of the analysis windows simultaneously.

To change the number of windows in the display

1. Tap the dropdown arrow for Rows and then select a different number.

   ![Rows Dropdown](image1)

2. Tap the dropdown arrow for Columns, and then select a different number.

   ![Columns Dropdown](image2)

You can also use the dropdown menus to change the windows that appear in the Dashboard.

To display a different window in a particular Dashboard slot

1. Tap the dropdown arrow in the upper-right corner to display the list of available windows, and then select a window.

   ![Window Dropdown](image3)

The dropdown list closes and the window changes immediately.
Dashboard Options

From the Dashboard window, depending on the individual dashboard window you have selected, you can use a range of options as described below:

- When you select a window, the options that appear in the menu bar correspond to the options you see if you open the same window from the side menu. For example, if you select the Dashboard Concentration chart, the Concentration menu options appear.

- You can reset the entire dashboard display by adding or removing rows or columns to display more or fewer individual windows.

  Tap or click Default View to return to the default display of two rows and two columns.

- From any window you can
  - Expand or collapse the window
  - Copy, save, or print the image
  - Select display, scale, axis, and logarithmic options
  - Use the dropdown list to replace the current display with a different display
  - Use the menu bar options for the particular chart type

When you add the Droplets chart to the Dashboard (highlighted below), QX Developer adds the View Mode options.

Note: The Droplets window is identical to the Event Counts window.

Table 18 on page 88 describes the option sets identified in the following graphic.
## Chapter 9 Data Analysis Module Overview

### Table 18. Dashboard options

<table>
<thead>
<tr>
<th>Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Border to identify selected window</td>
</tr>
<tr>
<td>2</td>
<td>Toolbar corresponding to the selected window type in the Dashboard</td>
</tr>
<tr>
<td>3</td>
<td>Dropdown arrow to replace the current display with a new display</td>
</tr>
<tr>
<td>4</td>
<td>Chart options</td>
</tr>
</tbody>
</table>
Plate View Windows

In the QX Developer Analysis module you can open the following windows that contain data on all wells in the run:

- **Plate Editor (analysis view)** — Displays the experiment type, sample names, sample types, supermixes, and assay type for each well in the run (similar to the Template Setup Plate Editor)
- **Plate View** — Displays concentration results in a text format for each processed well
- **2D Plate View** — Displays a two-dimensional amplitude chart for each well

**Plate Editor — Analysis View**

You can use the Plate Editor tab in the Analysis module to select wells, and then view or change the experiment parameters. Similar to other Plate Editor displays, the Analysis Plate Editor displays the experiment information for each well in the run.

*Important:* You can change all fields except the supermix.

Wells you select in the Analysis Plate Editor determine the data initially displayed in other analysis windows.
Plate View

The Plate View tab provides, within the plate layout grid, a text view of the calculated concentration for each target in each well processed in the run. For information on the concentration results in chart format, see Concentration on page 135.

<table>
<thead>
<tr>
<th>Well</th>
<th>Concentration (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>123.45</td>
</tr>
<tr>
<td>A2</td>
<td>234.56</td>
</tr>
<tr>
<td>A3</td>
<td>345.67</td>
</tr>
<tr>
<td>A4</td>
<td>456.78</td>
</tr>
<tr>
<td>A5</td>
<td>567.89</td>
</tr>
</tbody>
</table>

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2D Plate View

To facilitate comparisons of amplitude in the processed wells, the Plate 2D tab provides a two-dimensional amplitude view in a separate chart for each well. To see a larger representation of a well, select the well in the 2D Amplitude view.
Data Windows

QX Developer provides the following table views for your well and run data:

- **Data Table** — available from the main tabs on the left, shows your configured experiment parameters, plus all calculations from the run.

  *Note:* You can display the data in individual well format, merged format, or both.

- **Plate Well Data Table** — available in plate setup windows (including the Analysis module Plate Editor), shows only your configuration information for each individual well.

- **Analysis Well Data Table** — available in the Analysis module, shows a subset of configuration and run data for the wells selected in the Well Selector.

Data Table Tab

The Data Table window contains columns of data generated for each target in each well, as droplet clusters are identified through thresholding or manual clustering tools.
**Data Table Column Definitions**

Table 19 defines each field in the Data Table.

**Table 19. Data Table columns**

<table>
<thead>
<tr>
<th>Column name</th>
<th>Content description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Well</td>
<td>The well location in the plate</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> The instrument accepts 96-well plates only.</td>
</tr>
<tr>
<td>Sample description 1</td>
<td>Words or phrases to describe the sample in each well</td>
</tr>
<tr>
<td>Sample description 2</td>
<td><strong>Note:</strong> Only the sample description 1 data appears in plate layout views but you can pause on a well to show a tool tip containing the entries in all completed Sample description fields.</td>
</tr>
<tr>
<td>Sample description 3</td>
<td></td>
</tr>
<tr>
<td>Sample description 4</td>
<td></td>
</tr>
<tr>
<td>Target</td>
<td>Target name from Plate Editor tab</td>
</tr>
<tr>
<td>Conc(copies/µl)</td>
<td>Concentration of the target molecules recorded as copies per microliter</td>
</tr>
<tr>
<td>Molecular Weight (pg/µl)</td>
<td>Mass concentration of DNA in ddPCR well in pg/µl</td>
</tr>
<tr>
<td>Status</td>
<td>Contains either Check, OK, Multi, or Manual</td>
</tr>
<tr>
<td></td>
<td>• If Check, either the automatic analysis for the well failed or fewer than 10,000 droplets were counted in the well</td>
</tr>
<tr>
<td></td>
<td>• If OK, enough droplets were counted to trigger automatic analysis</td>
</tr>
<tr>
<td></td>
<td>• If Multi, the data was automatically analyzed as part of a multi-well selection</td>
</tr>
<tr>
<td></td>
<td>• If Manual, the droplets were analyzed manually</td>
</tr>
<tr>
<td>Status Reason</td>
<td>Identifies the reason for the status (unsuccessful automatic analysis, not enough droplets counted, or droplets analyzed manually)</td>
</tr>
<tr>
<td>Experiment</td>
<td>Experiment type selected for the well in the Plate Editor</td>
</tr>
<tr>
<td>SampleType</td>
<td>Sample type selected for the well in the Plate Editor</td>
</tr>
<tr>
<td>TargetType</td>
<td>Target type selected for each dye (FAM or EvaGreen® and HEX or VIC) in the Plate Editor</td>
</tr>
</tbody>
</table>
Table 19. Data Table columns, continued

<table>
<thead>
<tr>
<th>Column name</th>
<th>Content description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supermix</td>
<td>Supermix selected for the well in Plate Editor</td>
</tr>
<tr>
<td>DyeName(s)</td>
<td>Dye assigned to channels 1 and 2, respectively, for the well in the Plate Editor</td>
</tr>
<tr>
<td>Copies/20µlWell</td>
<td>Concentration of the target normalized to a volume of 20 µl</td>
</tr>
<tr>
<td>TotalConfMax</td>
<td>For merged wells the high error bar for the target concentration of the combined wells at a 95% confidence interval</td>
</tr>
<tr>
<td>TotalConfMin</td>
<td>For merged wells the low error bar for the target concentration of the combined wells at a 95% confidence interval</td>
</tr>
<tr>
<td>PoissonConfMax</td>
<td>Maximum target concentration normalized for the high error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>PoissonConfMin</td>
<td>Minimum target concentration normalized for the low error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>Accepted Droplets</td>
<td>Total number of droplets accepted by quality algorithm</td>
</tr>
<tr>
<td>Positives</td>
<td>Number of droplets that contain the target</td>
</tr>
<tr>
<td>Negatives</td>
<td>Number of droplets that do not contain the target</td>
</tr>
<tr>
<td>Ch1+Ch2+</td>
<td>Number of droplets that contain both channel 1 and channel 2 targets</td>
</tr>
<tr>
<td>Ch1+Ch2-</td>
<td>Number of droplets that contain just the channel 1 target</td>
</tr>
<tr>
<td>Ch1-Ch2+</td>
<td>Number of droplets that contain just the channel 2 target</td>
</tr>
<tr>
<td>Ch1-Ch2-</td>
<td>Number of droplets that contain neither channel 1 nor channel 2 targets</td>
</tr>
<tr>
<td>Linkage</td>
<td>Shows the concentration of copies of two targets that are predicted to be targeting a linked DNA molecule, calculated by an overabundance of copies in the double-positive cluster versus the expected value in copies per µl</td>
</tr>
<tr>
<td>CNV</td>
<td>Copy number calculated for the target relative to the reference</td>
</tr>
<tr>
<td>TotalCNVMax</td>
<td>For merged wells, the high error bar for the copy number of the combined wells at a 95% confidence interval</td>
</tr>
<tr>
<td>Column name</td>
<td>Content description</td>
</tr>
<tr>
<td>------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>TotalCNVMin</td>
<td>For merged wells, the low error bar for the copy number of the combined wells at a 95% confidence interval</td>
</tr>
<tr>
<td>PoissonCNVMax</td>
<td>Maximum copy number normalized for the high error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>PoissonCNVMin</td>
<td>Minimum copy number normalized for the low error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>ReferenceCopies</td>
<td>Copy number identified for the reference target in the Plate Editor Default is 2, indicating 2 copies per diploid genome.</td>
</tr>
<tr>
<td>UnknownCopies</td>
<td>This field is currently unused in QX Developer</td>
</tr>
<tr>
<td>Threshold1</td>
<td>Threshold value of first threshold line from left to right and bottom to top</td>
</tr>
<tr>
<td>Threshold2</td>
<td>Threshold value of second threshold line from left to right and bottom to top</td>
</tr>
<tr>
<td>Threshold3</td>
<td>Threshold value of third threshold line from left to right and bottom to top</td>
</tr>
<tr>
<td>ThresholdSigmaAbove</td>
<td>Distance of the threshold from the mean of the negative cluster, as a multiple of the negative cluster standard deviation</td>
</tr>
<tr>
<td>ThresholdSigmaBelow</td>
<td>Distance of the threshold from the mean of the positive cluster, as a multiple of the positive cluster standard deviation</td>
</tr>
<tr>
<td>ReferenceUsed</td>
<td>Identifies which target was used as a reference</td>
</tr>
<tr>
<td>Ratio</td>
<td>The ratio of the target against the identified reference</td>
</tr>
<tr>
<td>TotalRatioMax</td>
<td>For merged wells the high error bar for the ratio of the unknown against the reference of the combined wells at a 95% confidence interval</td>
</tr>
<tr>
<td>TotalRatioMin</td>
<td>For merged wells the low error bar for the ratio of the unknown against the reference of the combined wells at a 95% confidence interval</td>
</tr>
</tbody>
</table>
### Table 19. Data Table columns, continued

<table>
<thead>
<tr>
<th>Column name</th>
<th>Content description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PoissonRatioMax</td>
<td>Maximum ratio of the unknown against the reference normalized for the high error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>PoissonRatioMin</td>
<td>Minimum ratio of the unknown against the reference normalized for the low error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>FractionalAbundance</td>
<td>Calculation of fractional abundance of this unknown target vs. the reference target</td>
</tr>
<tr>
<td>TotalFractionalAbundanceMax</td>
<td>For merged wells the high error bar for the fractional abundance of the combined wells at a 95% confidence interval</td>
</tr>
<tr>
<td>TotalFractionalAbundanceMin</td>
<td>For merged wells the low error bar for the fractional abundance of the combined wells at a 95% confidence interval</td>
</tr>
<tr>
<td>PoissonFractionalAbundanceMax</td>
<td>Maximum fractional abundance normalized for the high error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>PoissonFractionalAbundanceMin</td>
<td>Minimum fractional abundance normalized for the low error bar of the droplet Poisson distribution for the 95% confidence interval</td>
</tr>
<tr>
<td>MeanAmplitudeOfPositives</td>
<td>Mean amplitude value of all droplets that contain the target</td>
</tr>
<tr>
<td>MeanAmplitudeOfNegatives</td>
<td>Mean amplitude value of all droplets that contain no target</td>
</tr>
<tr>
<td>MeanAmplitudeTotal</td>
<td>Mean amplitude value of all droplets</td>
</tr>
<tr>
<td>ExperimentComments</td>
<td>Not used in QX Developer.</td>
</tr>
<tr>
<td></td>
<td>Data populates this field only from files created in QuantaSoft versions 1.4 – 1.7, if comments were added.</td>
</tr>
<tr>
<td>MergedWells</td>
<td>Identifies which wells were merged together</td>
</tr>
<tr>
<td>TotalConfidenceMax68</td>
<td>For merged wells the high error bar for the target concentration of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>TotalConfidenceMin68</td>
<td>For merged wells the low error bar for the target concentration of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>PoissonConfidenceMax68</td>
<td>Maximum target concentration normalized for the high error bar of the droplet Poisson distribution for the 68% confidence interval</td>
</tr>
<tr>
<td>Column name</td>
<td>Content description</td>
</tr>
<tr>
<td>-------------------------------------</td>
<td>--------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>PoissonConfidenceMin68</td>
<td>Minimum target concentration normalized for the low error bar of the droplet Poisson distribution for the 68% confidence interval</td>
</tr>
<tr>
<td>TotalCNVMax68</td>
<td>For merged wells the high error bar for the copy number of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>TotalCNVMin68</td>
<td>For merged wells the low error bar for the copy number of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>PoissonCNVMax68</td>
<td>Maximum target concentration normalized for the high error bar of the droplet Poisson distribution for the 68% confidence interval</td>
</tr>
<tr>
<td>PoissonCNVMin68</td>
<td>Minimum target concentration normalized for the low error bar of the droplet Poisson distribution for the 68% confidence interval</td>
</tr>
<tr>
<td>TotalRatioMax68</td>
<td>For merged wells the high error bar for the ratio of the unknown against the reference of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>TotalRatioMin68</td>
<td>For merged wells the low error bar for the ratio of the unknown against the reference of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>PoissonRatioMax68</td>
<td>Maximum ratio of the unknown against the reference normalized for the high error bar of the droplet Poisson distribution for the 68% confidence interval</td>
</tr>
<tr>
<td>PoissonRatioMin68</td>
<td>Minimum ratio of the unknown against the reference normalized for the low error bar of the droplet Poisson distribution for the 68% confidence interval</td>
</tr>
<tr>
<td>TotalFractionalAbundanceMax68</td>
<td>For merged wells the high error bar for the fractional abundance of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>TotalFractionalAbundanceMin68</td>
<td>For merged wells the low error bar for the fractional abundance of the combined wells at a 68% confidence interval</td>
</tr>
<tr>
<td>PoissonFractionalAbundanceMax68</td>
<td>Maximum fractional abundance normalized for the high error bar of the droplet Poisson distribution for the 68% confidence interval</td>
</tr>
</tbody>
</table>
Table 19. Data Table columns, continued

<table>
<thead>
<tr>
<th>Column name</th>
<th>Content description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PoissonFractionalAbundanceMin68</td>
<td>Minimum fractional abundance normalized for the low error bar of the droplet Poisson distribution for the 68% confidence interval plate list after correcting the issue</td>
</tr>
<tr>
<td>TiltCorrected</td>
<td>Indicates if tilt correction was applied to the well</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Tilt correction rotates the angles from the double-negative cluster centroid to each droplet, such that the two single positive cluster centroids are orthogonal with the double-negative cluster centroid.</td>
</tr>
</tbody>
</table>
**Plate Well Data Table**

From the Plate Editor you can access a window containing the Well Data table, which displays configuration and run information on all wells containing sample.

**Note:** You cannot edit this table.

**Table 20. Plate Well Data Table columns**

<table>
<thead>
<tr>
<th>Column name</th>
<th>Content description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Well</td>
<td>Well number (A01 through H12)</td>
</tr>
</tbody>
</table>
| Perform Droplet Reading      | Yes, if wells are read  
No, if they are not | |
| Experiment Type              | Experiment Type selected in the Plate Editor:                                      |
|                              | - Direct Quantification (DQ)                                                        |
|                              | - Copy Number Variation (CNV)                                                       |
|                              | - Mutation Detection (MUT)                                                          |
|                              | - Rare Event Detection (RED)                                                        |
|                              | - Drop-off (DOF)                                                                   |
|                              | - Gene Expression (GEX)                                                            |
|                              | - Residual DNA Quantification (RDQ)                                                  |
| Sample description 1         | Words or phrases to identify each sample, as entered in the Plate Editor            |
| Sample description 2         |                                                                                     |
| Sample description 3         |                                                                                     |
| Sample description 4         |                                                                                     |

**Note:** You can enter up to four sample descriptions for each well.

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Sample type selected in the Plate Editor:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>- Unknown</td>
</tr>
<tr>
<td></td>
<td>- No template control (NTC)</td>
</tr>
<tr>
<td></td>
<td>- Positive control (Pos-Ctrl)</td>
</tr>
<tr>
<td></td>
<td>- Negative control (Neg-Ctrl)</td>
</tr>
</tbody>
</table>
### Table 20. Plate Well Data Table columns, continued

<table>
<thead>
<tr>
<th>Column name</th>
<th>Content description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supermix Name</td>
<td>Supermix selected in the Plate Editor:</td>
</tr>
<tr>
<td></td>
<td>- ddPCR Multiplex Supermix</td>
</tr>
<tr>
<td></td>
<td>- ddPCR Supermix for Probes (no dUTP)</td>
</tr>
<tr>
<td></td>
<td>- ddPCR Supermix for Probes</td>
</tr>
<tr>
<td></td>
<td>- ddPCR Supermix for Residual DNA Quantification</td>
</tr>
<tr>
<td></td>
<td>- One-Step RT-ddPCR Advanced Kit for Probes</td>
</tr>
<tr>
<td></td>
<td>- ddPCR EvaGreen® Supermix</td>
</tr>
<tr>
<td>Plex Mode</td>
<td>Assay method selected in the Plate Editor:</td>
</tr>
<tr>
<td></td>
<td>- Single Target per Channel</td>
</tr>
<tr>
<td></td>
<td>- Amplitude Multiplex</td>
</tr>
<tr>
<td></td>
<td>- Probe Mix Triplex</td>
</tr>
<tr>
<td></td>
<td>- Advanced Classification Method</td>
</tr>
<tr>
<td>Target Name</td>
<td>Target name entered in the Plate Editor tab</td>
</tr>
<tr>
<td>Target Type</td>
<td>Target type selected in the Plate Editor tab</td>
</tr>
<tr>
<td>Signal Channel 1</td>
<td>FAM or EvaGreen® fluorophore</td>
</tr>
<tr>
<td>Signal Channel 2</td>
<td>HEX or VIC fluorophore</td>
</tr>
<tr>
<td>Reference Copies</td>
<td>Copy number identified for the reference target in the Plate Editor tab</td>
</tr>
<tr>
<td>Well Notes</td>
<td>Entries by the user</td>
</tr>
<tr>
<td>Plot?</td>
<td>If selected, target will be used to calculate the copy number for the well</td>
</tr>
<tr>
<td>RDQ Conversion Factor</td>
<td>The Residual DNA Quantification conversion factor for the well</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> RDQ is only available in Residual DNA Quantification experiments.</td>
</tr>
</tbody>
</table>
Analysis Well Data Table

A well data table containing both configuration and run data appears in the analysis windows, next to the Well Selector. This table displays a subset of data columns for only the wells initially selected in the Plate Editor, or subsequently selected in the Well Selector.

Table 21 defines each column in the consolidated table.

Table 21. Fields in the Analysis Well Data table

<table>
<thead>
<tr>
<th>Column name</th>
<th>Content description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Well</td>
<td>The well location in the plate the sample is from</td>
</tr>
<tr>
<td>Sample description 1</td>
<td>The sample name used to identify the well in the Plate Editor tab</td>
</tr>
<tr>
<td>Sample description 2</td>
<td></td>
</tr>
<tr>
<td>Sample description 3</td>
<td></td>
</tr>
<tr>
<td>Sample description 4</td>
<td></td>
</tr>
<tr>
<td>Target</td>
<td>Target Name from Plate Editor tab</td>
</tr>
<tr>
<td>Conc (copies/µl)</td>
<td>Concentration identified using the Poisson calculation</td>
</tr>
<tr>
<td>Status</td>
<td>Notifies if clusters were identified or not and if they are the method employed (manual or auto analyze)</td>
</tr>
<tr>
<td>Experiment</td>
<td>Experiment type from the Plate Editor</td>
</tr>
<tr>
<td>SampleType</td>
<td>Sample type from the Plate Editor</td>
</tr>
<tr>
<td>TargetType</td>
<td>Target type assigned to the target in the Plate Editor</td>
</tr>
<tr>
<td>Supermix</td>
<td>Supermix from the Plate Editor</td>
</tr>
<tr>
<td>DyeName(s)</td>
<td>Fluorophore assigned to to a target in the Plate Editor</td>
</tr>
<tr>
<td>Column name</td>
<td>Content description</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------------------------------------------------</td>
</tr>
<tr>
<td>Accepted Droplets</td>
<td>Total number of droplets accepted by quality algorithms</td>
</tr>
<tr>
<td>Positives</td>
<td>Number of droplets that contain the target</td>
</tr>
<tr>
<td>Negatives</td>
<td>Number of droplets that do not contain the target</td>
</tr>
</tbody>
</table>
Analysis Windows

In its Analysis module, QX Developer provides amplitude displays and statistical probability charts in the Analysis module.

Amplitude Charts

Amplitude charts illustrate the droplets by degree of dye fluorescence. QX Developer displays fluorescence for positive and negative targets identified with FAM/EvaGreen® and HEX/VIC fluorophores in the following analysis windows:

- 1D Amplitude
- 2D Amplitude

Statistical Probability Charts

Statistical probability charts show calculations based on positive droplet concentrations. QX Developer counts the number of positive and negative droplets for each fluorophore in each sample, and then corrects the concentration based on a Poisson distribution to determine the concentration of the target DNA molecules in units of copies/µl in the following analysis windows:

- Concentration
- Copy Number
- Ratio
- Fractional Abundance

When you open the analysis data file in QX Developer and select one or more wells in the grid, the software plots each calculated Poisson probability distribution average as a data point in the grid. You can pause on the data point to see the high, median, and low numbers.

Note: You can display the data using a confidence level of either 68% or 95% (default setting).

Event Counts

Event counts record the number of droplets in which the target was found (positive) and was not found (negative). The Event Counts tab shows a bar chart view of the number of positive, negative, and total droplets for each of the targets in each of the selected wells.

Note: In the Dashboard display, event counts are shown in the Droplets window.

For more detailed information, see Chapter 10, Data Analysis Methodology.
Analysis Display and Output Options

QX Developer provides the following miscellaneous option types in the Analysis module:

- **Universal Options** — available from the primary toolbar to save, import, or export data files
- **Chart Display Preferences** — to change how your chart appears in the analysis window
- **Chart Scale Options** — to change the scale represented on an axis
- **Chart Menu Options** — available in chart views (from the icon) to copy, save, and print chart images. In Amplitude charts showing data points, you can show or hide
- **Table Menu Options** — available in table views (from the icon) to copy data, export data to Excel or CSV files, and show or hide columns

You can also expand the chart display to full-screen size and then return to the original display.

**Universal Options**

Universal options are explained in Table 22, and are available in all Analysis module windows.

**Table 22. Universal options**

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Save</td>
<td>Saves changes to the analysis file under its current file name or file extension</td>
</tr>
<tr>
<td>Save as</td>
<td>Saves the file as a different file extension (for example, .qlp to ddpcr to .ddpcr, or saves changes to the file under a different file name)</td>
</tr>
<tr>
<td>Import/export</td>
<td>Imports or exports plate setup data</td>
</tr>
<tr>
<td></td>
<td>Exports chart analysis data to an Excel file</td>
</tr>
<tr>
<td>Undo</td>
<td>Enabled in the Plate Editor only</td>
</tr>
<tr>
<td>User Settings</td>
<td>Allows you to restore your user preferences to the default settings and to show or hide color matrix change messages</td>
</tr>
</tbody>
</table>

**To save the data file**

1. Tap or click Save or Save As.
- If you tap or click Save, the software immediately saves any changes you made under the original file name.
- If you tap or click Save As, continue to Step 2.

2. In the Save As dialog box, enter a new file name, and then tap or click Save.

   The file is saved with the new file name.

To import or export setup or analysis data

1. Tap or click Import/Export.

   Table 23 describes the available import/export options.

   **Note:** If there is unsaved data, the software prompts you to save the file before its contents can be imported or exported.

**Table 23. Import/export options**

<table>
<thead>
<tr>
<th>Import/export option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Import plate setup CSV</td>
<td>Import plate layout data from a CSV file to the Plate Editor</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> This option is only available in the Plate Editor. The new plate layout overwrites the existing layout.</td>
</tr>
<tr>
<td>Export plate setup CSV</td>
<td>Export a configured Plate Editor layout to a CSV file</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> This option is only available in the Plate Editor.</td>
</tr>
<tr>
<td>Import plate setup</td>
<td>Import a plate layout from a plate template (ddplt) file</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> This option is only available in the Plate Editor.</td>
</tr>
<tr>
<td>Export plate setup</td>
<td>Export the plate layout to a plate template file.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> This option is only available in the Plate Editor.</td>
</tr>
<tr>
<td>Export visible data to CSV</td>
<td>Export only the data from the wells you selected in the Well Selector to a CSV file</td>
</tr>
<tr>
<td>Export cluster data</td>
<td>Exports the data to an Excel spreadsheet</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Includes (for each well) whether or not the target was found (1 = yes, 0 = no), the droplet count, and the mean and standard deviation for each dye, and the cluster ID</td>
</tr>
<tr>
<td>Export linkage data</td>
<td>Exports to a CSV file the number of copies of a target that are present in overabundance, versus. the expected value in copies per µl</td>
</tr>
</tbody>
</table>
Table 23. Import/export options, continued

<table>
<thead>
<tr>
<th>Import/export option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Export visible charts</td>
<td>Exports all charts that are displayed</td>
</tr>
<tr>
<td>Print visible charts</td>
<td>Prints all charts that are displayed</td>
</tr>
</tbody>
</table>

Table 24 shows where the options are available.

Table 24. Option availability

<table>
<thead>
<tr>
<th>Import/export option</th>
<th>Plate editor</th>
<th>Plate view</th>
<th>Plate 2D</th>
<th>Amplitude plots</th>
<th>Statistical probability charts</th>
<th>Event counts</th>
<th>Data table</th>
</tr>
</thead>
<tbody>
<tr>
<td>Import plate setup</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CSV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Export plate setup</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CSV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Import plate setup</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Export plate setup</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Export visible data</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>to CSV</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Export cluster data</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Export linkage data</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Export visible charts</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Print visible charts</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

Note: Amplitude plots are available in 1D and 2D, and statistical probability charts are Concentration, Copy Number, Ratio, and Fractional Abundance. In the Dashboard view corresponding options are available for each plot or chart displayed.
Expanding or Reducing the Chart

You can expand or reduce the chart size in the following windows:

- Dashboard
- 1D Amplitude
- 2D Amplitude
- Concentration
- Copy Number
- Ratio
- Fractional Abundance
- Event Counts

If you expand an amplitude or statistical probability chart to full size, the Well Selector and the Well Data table are hidden from view until you collapse the chart again.

To expand or reduce the chart size

1. To increase the chart size to the full window, tap or click the icon in the upper-right corner of the chart.
2. To reduce the window to its original size, tap or click the icon.
Selecting Chart Display Preferences

The Chart Display preferences primarily control font sizes and positioning. Table 25 explains each preference, as well as the windows in which it is available.

**Table 25. Chart display preferences**

<table>
<thead>
<tr>
<th>Preference</th>
<th>Window</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Channel or Dye</td>
<td>1D Amplitude 2D Amplitude</td>
<td>Select Channel or Dye to identify your chart accordingly. If you select Channel, the channel number appears as the chart title. If you select Dye, FAM or HEX appears as the chart title.</td>
</tr>
<tr>
<td>Axis Data Label Size</td>
<td>1D Amplitude 2D Amplitude</td>
<td>Select a different font size.</td>
</tr>
<tr>
<td></td>
<td>Concentration Copy Number Ratio Fractional Abundance Event Counts</td>
<td></td>
</tr>
<tr>
<td>Chart Data Label Size</td>
<td>Concentration Copy Number Ratio Fractional Abundance Event Counts</td>
<td>Select a different font size.</td>
</tr>
<tr>
<td>Chart Marker Size</td>
<td>Concentration Copy Number Ratio Fractional Abundance</td>
<td>Select a different font size.</td>
</tr>
<tr>
<td>Chart Marker Label Position</td>
<td>Concentration Copy Number Ratio Fractional Abundance</td>
<td>Reposition the marker label.</td>
</tr>
</tbody>
</table>
To select chart display preferences

1. Tap or click the Chart Display button.
2. Select your options.
   The display changes immediately to reflect your choices.
3. Do one of the following:
   - To save your choices as your default settings, tap or click Save as my default. When the success message appears, tap or click OK to close the dialog.
   - To close the menu without saving the settings, tap or click anywhere in the chart display.

Selecting Chart Scale Options

Chart scale options primarily control chart scale displays.

Each preference, as well as the windows in which it is available, is explained in Table 26.
Table 26. Chart scale options

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scale</td>
<td>Appears in options for all charts</td>
</tr>
<tr>
<td></td>
<td>The Scale option determines the numerical range for your chart data:</td>
</tr>
<tr>
<td></td>
<td>Auto — Selected by default; QX Developer automatically displays a scale to match the data distribution.</td>
</tr>
<tr>
<td></td>
<td>Fixed — Selecting Fixed enables the Axis fields, and allows you to manually adjust the range and increment.</td>
</tr>
<tr>
<td>Axis</td>
<td>Appears in options for all charts</td>
</tr>
<tr>
<td></td>
<td>When the Axis fields are enabled, you can manually adjust the minimum (Min) and maximum (Max) points for the data display on each axis, as well as the increments (Incr).</td>
</tr>
<tr>
<td></td>
<td>X — Horizontal axis</td>
</tr>
<tr>
<td></td>
<td>Y — Vertical axis</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> You can modify the x axis scale in the 2D Amplitude chart only.</td>
</tr>
<tr>
<td>Secondary Y Axis</td>
<td>Appears in options for Concentration only</td>
</tr>
<tr>
<td></td>
<td>From the dropdown list, you can select from Channel 2 concentration, ratio, copy number, or fractional abundance to display the corresponding data in a secondary row on the y axis.</td>
</tr>
<tr>
<td>Log</td>
<td>Appears in options for all statistical probability charts (Concentration, Copy Number, Ratio, and Fractional Abundance)</td>
</tr>
<tr>
<td></td>
<td>You can select the Log checkbox to display the y axis in base 10 logarithmic format.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> In the Concentration chart, you can also select Log for the secondary y axis.</td>
</tr>
<tr>
<td>Inverse</td>
<td>Appears in options for Ratio and Fractional Abundance only</td>
</tr>
<tr>
<td></td>
<td>The Inverse option equals the opposing number of the Ratio data point to a total of 2, and the opposing number of the Fractional Abundance data point to a total of 100.</td>
</tr>
<tr>
<td>Data labels</td>
<td>Appears in options for all statistical probability charts</td>
</tr>
<tr>
<td></td>
<td>A data label appears by default for each point in the chart, but you can clear the checkbox to remove the labels.</td>
</tr>
<tr>
<td>Grid Lines</td>
<td>Appears in options for all charts</td>
</tr>
<tr>
<td></td>
<td>Gridlines appear by default. You can clear the check box to remove them.</td>
</tr>
</tbody>
</table>
To select chart scale options

1. Tap or click the icon in the upper-left corner of the chart to display the corresponding chart options.

2. To manually define the chart scale, select Fixed.
   
   **Note:** Auto is the default setting.

3. In the Min, Max, and Inc fields, enter your lower and upper limits, and the increment, for the y axis. For 2D Amplitude, you can also change the scale on the x axis.

   **Note:** Increment entries are enabled only if the Log checkbox is cleared.

4. For Concentration, select an option for a secondary y axis display. To change to a base 10 logarithmic scale on the y axis, select the Log checkbox.

5. For Ratio and Fractional Abundance, to view an inverse calculation, select the Inverse checkbox.

6. To remove the data labels or grid lines from the display, clear the Data Labels or Grid Lines checkbox.
Using Chart Menu Options

Chart menu options are explained in Table 27.

Table 27. Chart menu options

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copy</td>
<td>Copies the chart as an image file, which you can paste into a separate document, such as an MS Word file</td>
</tr>
<tr>
<td>Save Image as</td>
<td>Saves the chart as an image file</td>
</tr>
<tr>
<td>Print</td>
<td>Prints the image</td>
</tr>
<tr>
<td>Show Point Values</td>
<td>Shows corresponding point values when you pause on a data point</td>
</tr>
</tbody>
</table>

Tip: Select the Show Point Values checkbox to see the data.
Chart menu options are available in the following windows:

- 1D Amplitude
- 2D Amplitude
- Concentration
- Copy Number
- Ratio
- Fractional Abundance
- Event Counts

**To use the chart menu options**

1. Select wells in the Well Selector.

Corresponding data appears in the Well Data table and the associated chart.
Chapter 9 Data Analysis Module Overview

2. In the upper-right corner of the chart, tap or click the menu icon (≡) and select from the dropdown list.
   - Select Copy to copy the chart to the Clipboard.
     
     **Note:** To paste the data, navigate to the location (for example, a Word file), then right-click and select Paste.
   - Select Save Image As and follow the prompts to save the file.
   - Select Print and follow the prompts to print the image.

3. Tap or click anywhere in the display to close the dialog box.

**Using Table Menu Options**

- Copy
- Export to Excel...
- Export to CSV...
- Show/Hide Columns

Table menu options are explained in Table 28.

**Table 28. Table options**

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copy</td>
<td>Copies the information from the selected wells</td>
</tr>
<tr>
<td>Export to Excel</td>
<td>Exports the data to an .xlsx file</td>
</tr>
<tr>
<td>Export to CSV</td>
<td>Exports the data to a .csv file</td>
</tr>
</tbody>
</table>
### Table 28. Table options, continued

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Show/Hide Columns</td>
<td>Add or remove columns from a table display</td>
</tr>
<tr>
<td></td>
<td>When you select Show/Hide Columns, the following dialog box opens.</td>
</tr>
<tr>
<td></td>
<td>Select or clear check boxes to show or hide columns in the data table.</td>
</tr>
</tbody>
</table>

- Select All
- Well
- Sample description 1
- Sample description 2
- Sample description 3
- Sample description 4
- Target
- Conc(copies/μL)
- Status
- Experiment
- SampleType
- TargetType
- Supermix
Table menu options are available in the following windows:

- 1D Amplitude
- 2D Amplitude
- Concentration
- Copy Number
- Ratio
- Fractional Abundance
- Event Counts

**Tip:** You can access the menu icon from the individual windows, or from the windows in the Dashboard display.

**To use the table menu options**

If you are using these options in the Data Table, skip to Step 3.

1. Select wells in the Well Selector.

   ![Well Selector](image)

   Corresponding data appears in the Well Data table.

   ![Well Data Table](image)
2. In the upper-right corner of the table, tap or click the menu icon (≡) and select from the dropdown list.
   - To copy the data from the selected wells to the Clipboard, select Copy.
     **Note:** To paste the data, navigate to the location (for example, an Excel file) and then right-click and select Paste.
   - To export your data, select Export to Excel or Export to CSV, and then follow the prompts to save the file.
     **Note:** Exporting to Excel exports the data in spreadsheet format, while Exporting to CSV exports the data in comma-delimited format.
   - To show or hide the corresponding column in the table, select Show/Hide Columns, and then select or clear checkboxes.

3. Tap or click anywhere in the display to close the dialog box.

**Viewing Run and Lot Information**

Use the Run Information window to view information on the run and view or add post-run notes regarding the plate.

For information on displaying your used lots in the Run Information window, see *Showing Lots in Run Information Output and Reports on page 62*. 
Chapter 9 Data Analysis Module Overview
Chapter 10 Data Analysis Methodology

As summarized in Chapter 9, Data Analysis Module Overview, the software presents analysis data in several different formats. When you open the data file in the Analysis module, you can view the data with the automatic analysis thresholds and calculations or you can choose different options to recalculate, replot, or recolor the results.

Data is shown in the following formats:

- **Amplitude**, which shows each droplet in each well in a scatter plot, in one-dimensional or two-dimensional views
- **Statistical models**, which show estimated target molecule concentration, ratio, and fractional abundance of one target against another target based on selected error model and confidence interval
  
  **Note:** Copy number variation is also calculated if the user ran a CNV experiment.
- **Event counts**, which are precise counts of positive and negative droplets for each target in each well

**Amplitude Plot Analysis Options**

QX Developer provides two amplitude plot displays in which you can view the data with automatic thresholding where applicable, or choose different options to replot or recolor the results.

In the 1D amplitude and 2D amplitude plots, you can

- Show well calculations
  
  **Note:** The 1D amplitude plot shows the data for each selected well in its own section, while the 2D amplitude plot shows all selected wells together.
- Modify thresholds
- View data in a heat map
- View histograms
- Re-apply automatic analysis
- Apply tilt correction to automatic analysis
Table 29 explains the analysis options that are available in both 1D amplitude and 2D amplitude plots.

### Table 29. 1D and 2D amplitude tool options

<table>
<thead>
<tr>
<th>Icon</th>
<th>Mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Lock Icon" /></td>
<td>View mode</td>
<td>Data appears with the thresholds or clusters assigned by QX Developer. When this icon is selected, the view is locked and you cannot modify the thresholds.</td>
</tr>
<tr>
<td><img src="image" alt="Heat Map Icon" /></td>
<td>Heat map</td>
<td>Shows the distribution of droplets, with higher density of droplets shown as hot.</td>
</tr>
<tr>
<td><img src="image" alt="Auto Icon" /></td>
<td>Automatic analysis</td>
<td>Disables tilt correction and automatically reanalyzes the data.</td>
</tr>
<tr>
<td><img src="image" alt="Auto with Tilt Icon" /></td>
<td>Automatic analysis with tilt correction</td>
<td>Enables tilt correction and automatically reanalyzes the data. <strong>Note:</strong> Tilt correction rotates the angles from the double-negative cluster centroid to each droplet, such that the two single positive cluster centroids are orthogonal with the double-negative cluster centroid.</td>
</tr>
<tr>
<td><img src="image" alt="Reset Icon" /></td>
<td>Reset</td>
<td>Resets the data to the original automatically analyzed data.</td>
</tr>
<tr>
<td><img src="image" alt="Chart Display Icon" /></td>
<td>Chart display options</td>
<td>Choose whether to display axis with channel number or fluorophore, and change the size of axis markers, data labels, and axis labels.</td>
</tr>
</tbody>
</table>
Table 30 explains the analysis options available only for 1D amplitude plots.

### Table 30. 1D amplitude tool options

<table>
<thead>
<tr>
<th>Icon</th>
<th>Mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Icon]</td>
<td>Threshold, single well</td>
<td>For a single well or multiple wells, the user can manually input thresholding values in the 1D amplitude plot.</td>
</tr>
<tr>
<td>![Icon]</td>
<td>Threshold, multiple wells</td>
<td>You can tap or click anywhere in the plot to apply threshold lines to the data.</td>
</tr>
<tr>
<td>![Icon]</td>
<td>Threshold, single well SD</td>
<td>For a single well or multiple wells, QX Developer automatically calculates and populates thresholding values with standard deviations (SD). <strong>Important:</strong> SD thresholds are not recommended for RMD, since these clusters are not typically located at right angles to each other. <strong>Note:</strong> The SD thresholds are set at a limit that is less than the mean number of positive targets, and more than the mean number of negative targets. Starting from where the user clicks in the plot, SD locates the threshold between the positive and negative clusters such that the classification error is minimized.</td>
</tr>
<tr>
<td>![Icon]</td>
<td>Threshold, multiple wells SD</td>
<td></td>
</tr>
<tr>
<td>![Icon]</td>
<td>Show histogram</td>
<td>Enables the icon to display your data in histogram plots</td>
</tr>
</tbody>
</table>
Table 31 explains the analysis options available only for 2D amplitude plots.

### Table 31. 2D amplitude tool options

<table>
<thead>
<tr>
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<th>Mode</th>
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</tr>
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</table>
| ![Threshold line icon](image) | Threshold line | Use the Threshold line icons to establish thresholds across a pair of channels, dividing droplets into distinct clusters separated by different colors. You can tap or click anywhere in the plot to apply threshold lines to the data. **To adjust the threshold values, you can**  
  - Drag the lines horizontally or vertically in the plot area  
  - Enter a new threshold value in the boxes at the end of each threshold line  
  - Tap the up and down arrows to incrementally adjust the numbers |
<p>| <img src="image" alt="Threshold line SD icon" /> | Threshold line SD | QX Developer automatically calculates and populates threshold line values with standard deviations (SD). <strong>Important</strong>: SD thresholds are not recommended for RMD, since these clusters are not typically located at right angles to each other. <strong>Note</strong>: The SD thresholds are set at a limit that is less than the mean number of positive targets, and more than the mean number of negative targets. Starting from where the user clicks in the plot, SD locates the threshold between the positive and negative clusters such that the classification error is minimized. |</p>
<table>
<thead>
<tr>
<th>Icon</th>
<th>Mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Threshold cluster square</td>
<td>Use the Threshold Cluster icons to establish thresholds across a pair of channels, dividing droplets into distinct clusters separated by different colors.</td>
</tr>
<tr>
<td></td>
<td>Threshold cluster circle</td>
<td>To classify your droplets, you can</td>
</tr>
<tr>
<td></td>
<td>Threshold cluster freeform</td>
<td>▪ Tap any of the three Threshold Cluster Mode buttons (square, circle, or free form)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>▪ Draw the chosen shape around the cluster of droplets</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Note:</strong> When using the cluster mode tools to draw a shape around a cluster of droplets, a dialog box with color selections by target appears in the window to help you select the correct target combination.</td>
</tr>
</tbody>
</table>

Target names are shown along the two axes based on the signal values assigned during plate setup. Tap or click the colored box that best represents the location of the cluster of droplets selected to automatically classify the droplets as positive or negative for each given target.

**Note:** The Cluster dialog box display varies according to the plex mode.
1D Amplitude

Use the 1D Amplitude analysis window to view droplet amplitudes for each channel, in acquisition order, for each well selected in the Well Selector.

The 1D Amplitude plots illustrate

- All accepted droplets, with a visual representation of the positive and negative droplets in each well

Positive droplets appear in color, while negative droplets appear in grayscale. The Well Selector displays an A in each well where QX Developer automatically thresholded the droplet data. If you create a manual threshold, the Well Selector displays an M in the wells where the threshold changed.
Amplitude in a histogram plot

The histogram displays frequency of droplets at specific fluorescence amplitudes for each well or group of wells selected. The Channel 1 (FAM/EvaGreen®) histogram appears in blue and Channel 2 (HEX/VIC) histogram in green.

Note: The Histogram icon is enabled by default so a Histogram plot appears automatically for each fluorophore (channel).

Amplitude in a Heat Map view

When you select the 1D Amplitude tab, the toolbar options that allow you to change thresholds and show or hide the heat map and histogram appear above the plots.
Chapter 10 Data Analysis Methodology

**Note:** For detailed information on each toolbar option, see Amplitude Plot Analysis Options on page 119.

Default color assignments for 1D amplitude plots are

- **Positive FAM** = Blue
- **Positive HEX** = Green
- **Negative** = Gray

The images in this section display data from a direct quantification experiment involving targets:

- The FAM plot (FAM dye in channel 1) shows the positive droplets for the first target in blue.
- The HEX plot (HEX dye in channel 2) shows the positive droplets for the second target in green.

Negative droplets appear in grayscale in each plot.

For experiments where enough droplets are recognized, QX Developer automatically configures an amplitude threshold, which distinguishes between positive and negative droplets.

**Note:** Well data must meet certain quality metrics (minimum of 10,000 droplets) before the software will automatically calculate a threshold above which droplets are considered positive. You can use the manual thresholding tools to ensure correct quantification of positive and negative droplets.
By default, the Histogram icon ( ) is enabled so the Histogram plot automatically appears with the amplitude plot.

To manually adjust the thresholds

1. Select one or more wells.

2. Tap or click any of the threshold icons ( ) to set manual thresholds:
   - Regular threshold for a single well; for multiple wells, you can set a different threshold for each well
   - Regular threshold for multiple wells, where you can set one threshold for all selected wells
   - Standard deviation threshold for a single well; for multiple wells, you can set a different threshold for each well
   - Standard deviation for multiple wells, where you can set one threshold for all selected wells

3. Tap or click on the location in the plot where you want to draw the new threshold.

4. When the line appears, use the up or down arrows, or click and drag the threshold, to adjust the location.
QX Developer displays the manual threshold and changes the color of the droplets that are now above the threshold.

**Note:** You can also manually enter a threshold number.

When you manually modify a threshold, the well selector displays an M for manual instead of the A for Automatic.

5. (Optional) To reset to the automatic threshold, tap or click Auto.

6. (Optional) To adjust the automatic threshold to consider tilt correction, tap or click Auto with Tilt.

7. (Optional) To reset everything to the original display, tap or click Reset.

**To view a heat map representation of the plot data**

▸ Tap or click the icon.
2D Amplitude

For the wells you select in the analysis view of the Plate Editor, the 2D Amplitude window illustrates fluorescence for two targets.

When you select the 2D Amplitude tab, the toolbar options allow you to change thresholds, manually cluster wells, and show or hide the heat map above the plots. For detailed information on each toolbar option, see Amplitude Plot Analysis Options on page 119.

Note: For detailed information on each toolbar option, see Amplitude Plot Analysis Options on page 119.

The images in this section display data from a direct quantification experiment involving two targets. The plot shows individual cluster representations for first and second target positive droplets, droplets containing both targets, and droplets containing neither target.
Default color assignments for 2D amplitude plots are

Positive FAM       =       Blue
Positive HEX/VIC   =       Green
Positive FAM and HEX/VIC = Orange (may appear brown on some monitors)
Double negative    =       Gray

Note: Well data must meet certain quality metrics (minimum of 10,000 droplets) before QX Developer will automatically calculate a threshold above which droplets are considered positive. You can use the manual thresholding tools to ensure correct quantification of positive and negative droplets.

To use the 2D amplitude tools

1. Open your data file in the Analysis module and select the 2D Amplitude tab.
2. Choose a threshold line or cluster option.

   Note: Use the cluster option when the data is not orthogonal and the threshold lines do not adequately divide the populations of droplets.

   - To adjust threshold lines, select a threshold line type and then tap or click in the plot to apply crosshair lines. Use any of the following methods:
     - Drag the lines horizontally or vertically in the plot area.
     - Enter a new threshold value in the boxes at the end of each threshold line.
     - Tap the up and down arrows to incrementally adjust the numbers.
To adjust thresholds using the cluster option, select a cluster type and draw the chosen shape around the cluster of droplets to classify.

3. (Optional) To reset to the automatic threshold, tap or click Auto.

4. (Optional) To adjust the automatic threshold to consider tilt correction, tap or click Auto with Tilt.

5. (Optional) To reset everything to the original display, tap or click Reset.
Statistical Probability Chart Analysis Options

QX Developer also provides chart displays where you can view the data from the original calculations, or choose different options to recalculate the results.

In charts reflecting statistical probability distributions, you can

- Show individual or merged well data.
- Change the error model and confidence level.

**Note:** For event counts, you can also alter the view mode to see positive, negative, or all droplet counts.
Table 32 explains the available options.

### Table 32. Chart tool options for statistical probability distribution charts

<table>
<thead>
<tr>
<th>Icon</th>
<th>Mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Both" /></td>
<td>Show both</td>
<td>Shows the calculated total or percentage as a merged calculation for all selected wells and as an individual calculation for each selected well</td>
</tr>
<tr>
<td><img src="image" alt="Merged" /></td>
<td>Show merged</td>
<td>Shows the calculated total or percentage as a combined number for all selected wells with the same parameters</td>
</tr>
<tr>
<td><img src="image" alt="Individual" /></td>
<td>Show individual</td>
<td>Shows the calculated total or percentage as an individual calculation for each selected well</td>
</tr>
<tr>
<td><img src="image" alt="Well" /></td>
<td>X-axis well</td>
<td>Shows the well number on the x axis</td>
</tr>
<tr>
<td><img src="image" alt="Sample" /></td>
<td>X-axis sample</td>
<td>Shows a partial sample name on the x axis, as well as the channel number</td>
</tr>
<tr>
<td><img src="image" alt="Error Model" /></td>
<td>Error model and confidence interval</td>
<td>Select an error model and confidence level. The default choices are Poisson and 95%, to yield the highest confidence in the results.</td>
</tr>
</tbody>
</table>

**Note:** Well setup must match for each well before the software can merge them.

**Important:** The software uses the following multipliers in its calculations:

- For 95% confidence interval, the sigma multiplier is 1.96.
- For 68% confidence interval, the sigma multiplier is 1.0.
Table 33 identifies the additional options available in the Event Count window. These options are available for event counts only.

Table 33. Additional options for event counts

<table>
<thead>
<tr>
<th>Icon</th>
<th>Mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td>View mode positive</td>
<td>Displays in a bar chart the number of positive droplets for each target in each well</td>
</tr>
<tr>
<td>Negative</td>
<td>View mode negative</td>
<td>Displays in a bar chart the number of negative droplets for each target in each well</td>
</tr>
<tr>
<td>Total</td>
<td>View mode total</td>
<td>Displays in a bar chart the number of total droplets in each well</td>
</tr>
</tbody>
</table>
Concentration measures the number of copies of the target molecules in each µl of sample based on the default settings. Use the toolbar to merge or separate the calculated results for each well, and to recalculate your results based on a different error model and confidence interval.

For information on toolbar options, see Statistical Probability Chart Analysis Options on page 132.

The following image displays concentration data, across a sequence of wells.

QX Developer plots the averaged concentration for each target as a data point. Pause on a data point to see the high, median, and low numbers.
The software uses the formula and variables shown below.

### Concentration Calculation

\[ c = -\ln \left( \frac{N_{\text{neg}}}{N} \right) / V_{\text{droplet}} \]

**Where:**

- \( \ln \): Negative natural logarithm
- \( V_{\text{droplet}} \): Volume of droplet
- \( N_{\text{neg}} \): Number of negative droplets
- \( N \): Total number of droplets

### Confidence Interval Calculation

**Note:** If you select a 68% confidence interval, the multiplier is changed from 1.96 to 1.00.

\[ P = \frac{N_{\text{neg}}}{N} \]

\[ \hat{\sigma}_{N_{\text{neg}}} = \sqrt{Np(1-p)} \]

\[ N_{\text{neg, \ max}} = N_{\text{neg}} + 1.96 * \hat{\sigma}_{N_{\text{neg}}} \]

\[ N_{\text{neg, \ min}} = N_{\text{neg}} - 1.96 * \hat{\sigma}_{N_{\text{neg}}} \]

**Where:**

- \( V_{\text{droplet}} \): Volume of droplet
- \( N_{\text{neg}} \): Number of negative droplets
- \( N \): Total number of droplets
- \( p \): p-value (probability of finding the target in the sample)
- \( \hat{\sigma} \): Delta
Bayesian Calculation

**Note:** If you specify the lesser confidence interval when you define your error model, the software uses 68% in its calculations instead of 95%.

\[
\begin{align*}
    b_1 &= -\ln\left(\frac{N_{\text{neg.min}}}{N}\right) / \text{V}_\text{droplet} \\
    b_2 &= -\ln\left(\frac{N_{\text{neg.max}}}{N}\right) / \text{V}_\text{droplet} \\
    (c_{\text{low, 95%}}, c_{\text{high, 95%}}) &= (\min(b_1, b_2), \max(b_1, b_2))
\end{align*}
\]

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<td>94</td>
<td>113.241</td>
<td>114.324</td>
</tr>
<tr>
<td>95</td>
<td>115.14</td>
<td>115.426</td>
</tr>
<tr>
<td>96</td>
<td>116.046</td>
<td>116.525</td>
</tr>
<tr>
<td>97</td>
<td>117.941</td>
<td>117.624</td>
</tr>
<tr>
<td>98</td>
<td>119.842</td>
<td>118.723</td>
</tr>
<tr>
<td>99</td>
<td>121.743</td>
<td>119.822</td>
</tr>
</tbody>
</table>
Copy Number

**Important:** To see copy number variation data, you must run a copy number variation (CNV) ddPCR experiment.

Copy number measures the probable count of the target gene, relative to the reference gene, to calculate the number gene targets per genome. Use the toolbar to merge or separate the calculated results for each well, and to recalculate your results based on a different error model and confidence interval. For information on toolbar options, see Statistical Probability Chart Analysis Options on page 132.

The following image displays copy number data, across a sequence of wells.

QX Developer plots each copy number variation as a data point. Pause on a data point to see the high, median, and low numbers.
The software uses the formula and variables shown below.

\[ CNV = \frac{A}{B} N_B \]

Where:

<table>
<thead>
<tr>
<th></th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Target concentration</td>
</tr>
<tr>
<td>B</td>
<td>Reference concentration</td>
</tr>
<tr>
<td>N_B</td>
<td>Number of reference copies in genome (usually 2)</td>
</tr>
</tbody>
</table>

### Confidence Interval Calculations

**Note:** If you specify the lesser confidence interval when you define your error model, the software uses 68% in its calculations instead of 95%.

\[ CI_{CNV} = N_B \cdot \frac{A}{B} \sqrt{\frac{CI_A^2}{A^2} + \frac{CI_B^2}{B^2}} \]

\[
(CNV_{low\_95\%}, CNV_{hi\_95\%}) = (CNV - (1/2)CI_{CNV}, CNV + (1/2)CI_{CNV})
\]

Where:

<table>
<thead>
<tr>
<th></th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Target concentration</td>
</tr>
<tr>
<td>B</td>
<td>Reference concentration</td>
</tr>
<tr>
<td>N_B</td>
<td>Number of reference copies in genome (usually 2)</td>
</tr>
<tr>
<td>CI_A</td>
<td>Poisson confidence interval of target (95% max - 95% min)</td>
</tr>
<tr>
<td>CI_B</td>
<td>Poisson confidence interval of reference (95% max - 95% min)</td>
</tr>
</tbody>
</table>

**Notes:**

Copy Number analysis involves determining the number of copies of a given target DNA sequence as compared to an invariant reference DNA sequence. Use the CNV analysis to determine structural variations, such as ploidy changes (addition or deletion of chromosomes), deletions (missing partial chromosomes) or duplications (repeated target sequences) in the genome.
The ability to discriminate, with statistical confidence, between consecutive CN states has been a technical challenge in copy number assessment. Fundamentally, as CN state increases, the percentage difference in target genomic material between states decreases. For example, for a given target locus, a CN of 3 is 50% more abundant in concentration per genome than a CN of 2, while a CN of 5 is only 25% more abundant per genome than a CN of 4, and so on.

Due to the massive partitioning of the reaction across up to 20,000 droplets per well, and the absolute nature and precision of the ddPCR concentration measurement, ddPCR CNV analysis enables the quantitative discrimination required to resolve consecutive copy number states beyond CN 3. For example, a 20% difference in target concentration between a CN 5 versus CN 6 sample can be reproducibly resolved using ddPCR. This discrimination is both a function of concentration differences between consecutive CN states, as well as the amount of DNA assayed per well.
Ratio

Ratio determines the quotient of two target molecules in each µl of sample based on the default settings.

T1:T2

Use the toolbar to merge or separate the calculated results for each well, and to recalculate your results based on a different error model and confidence interval. For information on toolbar options, see Statistical Probability Chart Analysis Options on page 132.

The following image displays ratio data, across a sequence of wells.

QX Developer plots each ratio as a data point. To see the confidence level numbers, pause on the data point.
Chapter 10 Data Analysis Methodology

The software uses the formula and variables shown below.

\[ r = \frac{A}{B} \]

**Where:**

- **A**  Target concentration
- **B**  Reference concentration

**Note:** If no reference is selected, the highest channel will automatically be considered as the reference. If multiple references are selected, the geometric mean is used.

**Confidence Interval Calculations**

**Note:** If you specify the lesser confidence interval when you define your error model, the software uses 68% in its calculations instead of 95%.

\[ CI_r = \frac{A}{B} \sqrt{\frac{CI_A^2}{A^2} + \frac{CI_B^2}{B^2}} \]

\[ (r_{low\_95\%}, r_{hi\_95\%}) = (r - (1/2)CI_r, r + (1/2)CI_r) \]

**Where:**

- **A**  Channel 1 target concentration
- **B**  Channel 2 target concentration
- **CI_A**  Confidence interval of Channel 1 target (95% max - 95% min)
- **CI_B**  Confidence interval of Channel 2 target (95% max - 95% min)
**Fractional Abundance**

Fractional Abundance determines the relative abundance of your targets in each µl of sample based on the default settings. Use the toolbar to merge or separate the calculated results for each well, and to recalculate your results based on a different error model and confidence interval. For information on toolbar options, see *Statistical Probability Chart Analysis Options on page 132.*

The following image displays fractional abundance data across a sequence of wells.

QX Developer plots the fractional abundance as a data point. To see the high and low confidence interval numbers, pause on the data point.

The software uses the formula and variables shown below.

\[ f = \frac{A}{A + B} \]
Chapter 10 Data Analysis Methodology

Where:

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Target concentration</td>
</tr>
<tr>
<td>B</td>
<td>Reference concentration</td>
</tr>
</tbody>
</table>

**Note:** If no reference is selected, the highest channel will automatically be considered as the reference. If multiple references are selected, the geometric mean is used.

**Confidence Interval Calculations**

**Note:** If you specify the lesser confidence interval when you define your error model, the software uses 68% in its calculations instead of 95%.

\[
CI_f = \frac{1}{(A + B)^2} \sqrt{B^2CI_A^2 + A^2CI_B^2}
\]

\[
(f_{low\_95\%}, f_{hi\_95\%}) = (f - (1/2)CI_f, f + (1/2)CI_f)
\]

Where:

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Channel 1 target concentration</td>
</tr>
<tr>
<td>B</td>
<td>Channel 2 target concentration</td>
</tr>
<tr>
<td>CI_A</td>
<td>Confidence interval of Channel 1 target (95% max - 95% min)</td>
</tr>
<tr>
<td>CI_B</td>
<td>Confidence interval of Channel 2 target (95% max - 95% min)</td>
</tr>
</tbody>
</table>
The Event Counts tab shows a bar chart view of the number of positive and negative droplets for each of the targets in each of the selected wells.

Note: In the Dashboard display, event counts appear in the Droplets window.

Use the toolbar to change the display by combining or separating data for each well. You can also show data by well name or by sample name, and you can show any combination of positive, negative, or combined droplet counts.
Linkage Analysis

In ddPCR, physical linkage of two targets can occur in one of the following patterns;

- Tandem repeats if the same target sequence
- Proximity of two target sequences physically linked on the same piece of DNA

QX Developer uses the formula and variables shown below.

\[
 linkage = \ln(N) - \ln\left(n_{00} + n_{01} + n_{10} + \frac{n_{01}n_{10}}{n_{00}}\right)
\]

Where:

- \(\ln\) Natural logarithm
- \(-\ln\) Negative natural logarithm
- \(N\) Total number of droplets
- \(n_{00}\) Number of negative droplets for both channels
- \(n_{01}\) Number of negative droplets in channel 1 and positive droplets in channel 2
- \(n_{10}\) Number of positive droplets in channel 1 and negative droplets in channel 2

\[
\sigma_{\text{linkage}} = \sqrt{\frac{(n_{00} + n_{01})^2 \sigma_{10}^2 + (n_{00} + n_{10})^2 \sigma_{01}^2 + (n_{00} - \frac{n_{10}n_{01}}{n_{00}})^2 \sigma_{00}^2}{(n_{00} + n_{01} + n_{01}n_{10} + n_{00}n_{01} + n_{01}n_{10})^2}}
\]

Where:

- \(N\) Total number of droplets
- \(n_{00}\) Number of negative droplets for both channels
- \(n_{01}\) Number of negative droplets in channel 1 and positive droplets in channel 2
- \(n_{10}\) Number of positive droplets in channel 1 and negative droplets in channel 2
\( p \)  \( \text{p-value (calculated probability)} \)

\( \delta \)  \( \text{Delta} \)

**Confidence Interval Calculation**

\[
CI_{\text{linkage}} = (\text{linkage} - (1.96)\sigma_{\text{linkage}}, \text{linkage} + (1.96)\sigma_{\text{linkage}})
\]

Where:

\( \sigma \)  \( \text{Standard deviation} \)
Chapter 11 Analysis Module Reporting

Use the Reporting function to create a variety of reports in the QX Developer Analysis module.

You can create reports on the fly from any analysis file or, if you need to generate specific reports on a regular basis, you can set up report templates for reuse.

This section contains information on available report elements, how to create analysis reports, and how to create report templates.

Report Elements

QX Developer allows you to generate reports for any or all of the following elements of the run and data.

Run Setup

- Plate Setup
  - Well
  - Perform droplet reading
  - Experiment type
  - Sample description 1, 2, 3, and 4
  - Sample type
  - Supermix name
  - Plex mode
  - Target name
  - Target type
  - Signal channel 1 and 2
  - Reference copies
  - Well notes
  - Plot?
Rdq conversion factor

- Plate name
- Supermix information
- Number of wells read
- Selected wells for report
- Username of the person who did the run
- Run start time
- Run complete time
- Initial wait elapsed time
- DR elapsed time
- Errors
- Instrument serial number
- Firmware version
- Software version
- Software
- Run notes
- Lot information

1D Amplitude

- Threshold value
- Tilt correction
- Amplitude 1D chart
- Amplitude 1D chart for each well
- Amplitude 1D histogram chart
- Amplitude 1D histogram chart for each well

2D Amplitude

- Threshold value
- Tilt correction
Amplitude 2D chart
Amplitude 2D chart for each well

**Concentration**
- Error model
- Confidence model
- Chart

**Copy Number**
- Error model
- Confidence model
- Chart

**Ratio**
- Error model
- Confidence model
- Inverse
- Chart

**Fractional Abundance**
- Error model
- Confidence model
- Inverse
- Chart

**Event Counts**

**Analysis Results**

**Audit Logs**

The audit log contains all of the original data from the plate run, and also records any changes that were made to the data file. The log records the change number, date, username for the user who made the change, the software version, full user name, computer name, and the reason for the change. Also recorded in a tabular format are the change type and description, as well as old and new
values for all well settings. The description column includes the well number of any affected well, and may also include other descriptive information.
Creating Analysis Reports

You can create a report using a template or you can create a custom report on the fly.

1. Select the Reports tab.
2. Do one of the following:
   - To create a report from a template, click the Template drop-down arrow and select a template from the list.
   - To change reporting elements in the template, go to Step 3.
   - To run the report without changing the template, go to Step 5.
   - To create a custom report, go to Step 3.
3. Click the arrow on the left to see expanded lists.
4. Select checkboxes for report elements to include.
5. Enter a report title.
6. (Optional) Select the Password Encrypt Report checkbox and then enter a password.
7. Click Generate Report.
   - The software creates a PDF report.
8. To save the report, click Save.
Creating Report Templates

If you are assigned the Create New Templates user privilege, you can create and save a new report template by editing the default report structure or by opening and editing an existing report, and then saving the modified structure as a new template.

Note: For information on creating reports, see Report Elements on page 149.

To create a report template

1. Select the Template Setup tab and tap or click Reports.

2. Do one of the following:
   - To create a new template, tap or click Add New Template.
   - To open an existing template, select the template and then tap or click Edit Report Template.
   The default or selected layout appears.

3. Tap or click the arrow to the left of each item to expand the list of reportable items.
4. Select and clear checkboxes to create the report structure.

   **Tip:** Use the buttons at the bottom of the screen to expand or collapse all categories at once, or to select or clear all checkboxes at once.

5. In the Report Title field, enter a name for your report template. You cannot exceed 30 characters.

6. Tap or click Save.

7. In the Save dialog box, enter a name for your template file.

8. Select a storage location.

   Choose from the following:

   - System Templates or My Templates

   **Important:** Only one of these locations will be valid. If your administrator has specified preferred Locations in System Settings, you can save your templates to the System
Chapter 11 Analysis Module Reporting

Templates folder or the Shared folder. If preferred locations are not specified, you can save them to the My Templates folder or the Shared Templates folder.

- **Shared Templates**

  You have the option to save your templates to this folder to make them available to all users.

9. Tap or click Save again to save the report format as a template.
Chapter 12 Gene Study Module

You access the Gene Study module from the Data Analysis window.

To create a gene study, you add one or more data files and then compare gene expression data from one or more ddPCR experiments.

Note: The maximum number of samples you can analyze in a gene study is limited by the size of the computer's RAM and virtual memory.

To open the Gene Study module

► Click Gene Study.
The Gene Study dialog box includes two tabs:

- **Study Setup** — manages the run files in the gene study
  
  *Important:* Adding or removing data files in a gene study does not change the data in the original file.

- **Study Analysis** — displays the gene expression data for the combined runs.

From these tabs you can add files to set up a gene study, analyze your data in charts and plots, and run reports on the gene study data.

### Gene Study Options

The Gene Study module provides the menus and corresponding options described below.

- **File** — Open, save, and close the gene study, and add files

- **Settings** — Restore the default layout

- **Tools** — Create a report
To open a gene study file

1. Click File ➔ Open.
2. In the Open dialog box, navigate to and select the data file, and then tap or click Open.
   The file opens in the Gene Study module.

To save a gene study file

1. Click File and select Save or Save As.
2. If prompted, enter a file name and tap or click Save.
3. If prompted to overwrite an existing file, tap or click Yes or No.

To close a gene study file

► Click File ➔ Close.
   Note: If you have not saved the file, an advisory prompt appears.

To add data files

1. Click File ➔ Add Data Files.
2. In the Open dialog box, navigate to the data files to be added to the study.
3. Select one or more files and then tap or click Open.
   The files are added to the Study Setup window.

To restore default settings

► Click Settings ➔ Restore default settings.
   Note: At least one modified data file must be open.

To generate a report

► See Gene Study Reports on page 167.
Setting up the Gene Study

Use the Study Setup window to

- Add data files to include in the study
- Remove data files from the study
- Select data files for viewing and analysis
- Show the plate view for a data file

Table 34 defines each column in Study Setup.
### Table 34. Gene Study Setup Fields

<table>
<thead>
<tr>
<th>Column Title</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>File Name</td>
<td>Name of the data file</td>
</tr>
<tr>
<td>Date Created</td>
<td>Date the file was imported into the Gene Study module</td>
</tr>
<tr>
<td>Plate View</td>
<td>Opens a map of the plate with the data for each dye in each well included in the Gene Study</td>
</tr>
</tbody>
</table>
Adding Data Files

To add data files

1. On the Study Setup tab, tap or click Add Data Files.
2. In the Open dialog box, navigate to the data files to be added to the study.
3. Select one or more files and then tap or click Open.

The files are added to the Study Setup window.

Removing Data Files

To remove data files

- Select the checkbox next to the file and then tap or click Remove.

The file is removed from the Study Setup window.

Note: The Remove button is disabled until you select a file.
Selecting Data Files for Analysis

To select data files for analysis

1. Select the checkboxes next to the files to be included in the analysis.

2. Select the Study Analysis tab to see the analysis results. For information, see Using the Study Analysis Features on page 165.

Showing the Plate View

To show the plate view

1. In the Plate View column, tap or click the button for the plate view to be displayed.

The Plate view appears.
2. Click Close to close the view.
Using the Study Analysis Features

The Study Analysis tab displays the data from all runs in the gene study in the views described in Table 35.

Table 35. Analysis Toolbar

<table>
<thead>
<tr>
<th>Button</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bar Chart</td>
<td>Size of bars correspond to normalized expression or relative quantity</td>
</tr>
<tr>
<td>Clustergram</td>
<td>Data is arranged in a hierarchy based on the degree of similarity of expression for different targets and samples</td>
</tr>
</tbody>
</table>
Table 35. Analysis Toolbar, continued

<table>
<thead>
<tr>
<th>Button</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scatter Plot</td>
<td>Dot plot of normalized expression of targets for a control versus an experimental sample</td>
</tr>
<tr>
<td>Volcano Plot</td>
<td>Dot plot of change in expression (regulation) of a target for an experimental sample compared to a control</td>
</tr>
<tr>
<td>Heat Map</td>
<td>Plate layout grid showing target regulation for an experimental compared to a control sample based on relative normalized expression</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> If the same targets are in the same location on multiple plates but with different samples, use the dropdown menu to select a particular plate for analysis.</td>
</tr>
<tr>
<td>Results Table</td>
<td>Provides results in a tabular format</td>
</tr>
</tbody>
</table>
Gene Study Reports

Use the Gene Study Report dialog box to arrange the gene study data into a report. Table 34 lists all the options available for a gene study report.

Table 36. Gene Study Report Categories

<table>
<thead>
<tr>
<th>Category Header</th>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Report Information</td>
<td>Date, user name, data file name, data file path, and the selected well group</td>
</tr>
<tr>
<td></td>
<td>Gene Study File List</td>
<td>List of all the data files in the gene study</td>
</tr>
<tr>
<td></td>
<td>Notes</td>
<td>Notes about the data report</td>
</tr>
<tr>
<td>Study Analysis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bar Chart</td>
<td>Analysis Settings</td>
<td>List of the selected parameters</td>
</tr>
<tr>
<td></td>
<td>Chart</td>
<td>Gene expression bar chart showing the data</td>
</tr>
<tr>
<td></td>
<td>Target Names</td>
<td>List of targets in the gene study</td>
</tr>
<tr>
<td></td>
<td>Sample Names</td>
<td>List of samples in the gene study</td>
</tr>
<tr>
<td></td>
<td>Data</td>
<td>Spreadsheet that shows the data</td>
</tr>
<tr>
<td></td>
<td>Target Stability</td>
<td>Target stability data</td>
</tr>
<tr>
<td></td>
<td>Inter-run Calibration</td>
<td>Inter-run calibration data</td>
</tr>
<tr>
<td>Study Analysis</td>
<td>Clustergram, Scatter Plot, Volcano Plot, and Heat Map</td>
<td></td>
</tr>
<tr>
<td>Bar Chart</td>
<td>Analysis Settings</td>
<td>Settings for each chart type</td>
</tr>
<tr>
<td></td>
<td>Chart</td>
<td>Gene expression chart showing the data</td>
</tr>
<tr>
<td></td>
<td>Data</td>
<td>Spreadsheet listing the data in each target</td>
</tr>
</tbody>
</table>
Creating a Gene Study Report

Before creating a report, make any needed adjustments to the gene study data and charts.

To create a gene study report

1. Select Tools ➔ Reports in the Gene Study menu to open the Report dialog box.
2. Select the options you want to include in the report.
   - The report opens with default options selected. Select or clear the checkboxes to change whole categories or individual options within a category.
   - For information on the available options to display, see Gene Study Reports on page 167.
3. Change the order of categories and items in a report.
   - Drag the options to the required position. Note that items can be reordered only within the categories to which they belong.
4. Click Update Report to update the Report Preview window with any changes.
5. Print or save the report:
   a. Click the Print Report button in the toolbar to print the current report.
   b. Select File ➔ Save to save the report in PDF, MHT or MHTML file format and select a location to save the file.
      - Note: MHT and MHTML are Microsoft formats.
   c. Select File ➔ Save As to save the report with a new name or in a new location.
6. (Optional) To save the current report settings as a template, select Template ➔ Save or Save As.
Appendix A Experiment Examples

This section contains setup and analysis information for the following sample experiments:

- Direct Quantification, using the Amplitude Multiplex assay type
- Copy Number Variation, using the Probe Mix Triplex assay type
- Drop-Off, using the Basic Drop-off Assay type for genome edit detection
- Direct Quantification, using the Advanced Classification Method assay type

Important: This appendix is intended for use with data acquisition files where the supermix and wells have been selected and cannot be changed during analysis.
Direct Quantification Multiplexed

You can increase the number of targets to be evaluated by using the Amplitude Multiplex option.

Setting Up Experiment Parameters

To set experiment parameters

1. Open the analysis data file and select the Plate Editor tab.

2. Select one or more wells to be assigned well analysis attributes.

3. Change the experiment type to Direct Quantification (DQ), and then tap or click Apply.

4. (Optional) Change the sample description identifiers.

5. (Optional) Change the sample type. The default type is Unknown.
   
   Note: You cannot change the supermix that was used during data collection.

6. Under Assay Type, select Amplitude Multiplex and tap or click Apply.
   
   Four target rows appear.

7. Change the target name and type (unknown or reference) for each target in the well. The software defaults to FAM Lo, FAM Hi, HEX Lo, and HEX Hi, respectively, for each target.

8. If you select Reference, you can change the number of reference copies. The default is 2.
   
   If no reference is selected, the target in the highest channel is used as the reference.
If more than one reference is selected in the well, checkboxes appear to the right for using the reference in the plot.

**Note:** Each Reference target enables ratio calculations in the results.

9. (Optional) Change the high and low target identifiers.

Amplitude Multiplex allows for up to two assays in each channel, indicated by Lo and Hi labels. In this multiplex mode, each target may have a signal for only one channel. You can

- Select a different dye, if the channel supports two dyes (for example, change HEX to VIC or VIC to HEX)
- Select a different channel for the target (for example, change FAM to HEX)
- Change Hi to Lo or Lo to Hi

10. Tap or click Apply.

**Tip:** Optionally, to expedite plate setup you can copy and paste data from one well to another, and you can tap or click Undo to undo an applied well setting.

**Viewing and Adjusting Results**

Automatic thresholding is not available for Amplitude Multiplex experiments.

For clusters that are easily identifiable across the chart, use Threshold Lines mode to set manual thresholds. If the data is not orthogonal and the threshold lines do not adequately divide the populations of droplets, use Threshold Clusters mode to set manual thresholds.

**To set thresholds in the 2D Amplitude window**

1. Select the 2D Amplitude tab.

2. On the toolbar, tap or click the Threshold Line Mode button.
3. Tap or click anywhere on the plot to apply “best fit” threshold lines to the data.

4. To adjust the thresholds, drag the lines horizontally or vertically in the plot area.

**Tip:** You can also enter a specific threshold value in the boxes at the end of each threshold line.

![Plot with threshold lines](image)

**To apply and adjust manual clusters**

1. Tap or click any of the three cluster mode buttons (square, circle, or freehand).

2. Press the left mouse button and draw the chosen shape around the cluster of droplets to be classified first.

**Tip:** When you release the mouse button, a pop-up tool appears to help you select the correct target combination. Target names are shown along the two axis based on the signal values assigned during plate setup.

![Select to assign cluster](image)

3. To classify droplets as positive or negative for each given target, select the colored box that best represents the location of the cluster of droplets.
Copy Number Variation Triplexed

A CNV experiment with probe mix triplexing uses two targets and a reference to produce analysis data.

Setting Up Experiment Parameters

To set experiment parameters

1. Open the analysis data file and select the Plate Editor tab.
2. Select one or more wells to be assigned well analysis attributes.
3. Set the experiment type to Copy Number Variation (CNV), and tap or click Apply.
4. (Optional) Change the sample description identifiers.
5. (Optional) Change the sample type. The default type is Unknown.
   
   **Note:** You cannot change the supermix that was used during data collection.

6. Tap or click Apply.
7. From the Assay Type dropdown list, select Probe Mix Triplex and tap or click Apply.

   QX Developer displays three targets.

   **Important:** You cannot add or delete triplex targets.

8. Enter target names and select the target type (unknown or reference) for each target in the well.

9. Assign the targets as follows:

<table>
<thead>
<tr>
<th>Target</th>
<th>Target Type</th>
<th>Signal Ch1</th>
<th>Signal Ch2</th>
<th>Ref Copies</th>
<th>Plot?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target 1</td>
<td>Unknown</td>
<td>FAM</td>
<td>None</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Target 2</td>
<td>Unknown</td>
<td>None</td>
<td>HEX/VIC</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Target 3</td>
<td>Reference</td>
<td>FAM</td>
<td>HEX/VIC</td>
<td>2</td>
<td>Yes</td>
</tr>
</tbody>
</table>

10. Tap or click Apply.

**Tip:** Optionally, to expedite plate setup you can copy and paste data from one well to another, and you can tap or click Undo to undo an applied well setting.
Viewing and Adjusting Results

**Note:** Thresholding is not available for Probe Mix Triplex experiments.

**To see the results**

1. After all wells are configured, select the 2D Amplitude tab to evaluate the data and manually set clusters.
2. Tap or click any of the three cluster mode buttons (square, circle, or freehand).
3. Press the left mouse button and draw the chosen shape around the cluster of droplets to be classified first.
   **Tip:** When you release the mouse button, a pop-up tool appears to help you select the correct target combination. Target names are shown along the two axis based on the signal values assigned during plate setup.

4. To classify droplets as positive or negative for each given target, select the colored box that best represents the location of the cluster of droplets.
Drop-Off Assays for Genome Edit Detection

Two configurations of drop-off assays are possible and are shown below. Assign a signal (expected cluster position) to wild-type and NHEJ indel alleles. The signal produced by a wild-type allele will always be positive for both FAM and HEX, while the signal produced by an NHEJ indel allele depends on the assay configuration used (as shown below left and right).

**Drop-Off Probe in HEX**

\[ \text{NHEJ Probe} = \text{HEX and Reference Probe} = \text{FAM} \]

**Drop-Off Probe in FAM**

\[ \text{NHEJ Probe} = \text{FAM and Reference Probe} = \text{HEX} \]

Setting Up Experiment Parameters

To set experiment parameters

1. Open the analysis data file and select the Plate Editor tab.
2. Select one or more wells to be assigned well analysis attributes.
3. Set the experiment type to Drop-Off (DOF) and tap or click Apply.
4. (Optional) Change the sample description identifiers.

5. (Optional) Change the sample type. The default type is Unknown.

    **Note:** You cannot change the supermix that was used during data collection.

6. Tap or click Apply.

    QX Developer automatically assigns the assay method as Basic Drop-Off.

7. Enter the target names, and then designate one target as Reference and the other as Unknown.

    The Reference target enables ratio calculation of results.

8. Tap or click Apply.

    **Tip:** Optionally, to expedite plate setup you can copy and paste data from one well to another, and you can tap or click Undo to undo an applied well setting.

### Viewing and Adjusting Results

Thresholding is not available for Drop Off experiments.

**To apply and adjust clusters**

1. After all wells are configured, select the 2D Amplitude tab to evaluate the data and manually set clusters.

2. Tap or click any of the three cluster mode buttons (square, circle, or freehand).
3. Press the left mouse button and draw the chosen shape around the cluster of droplets to be classified first.

**Tip:** When you release the mouse button, a pop-up tool appears to help you select the available cluster assignments. Target names are shown along the two axis based on the signal values assigned during plate setup.

4. To classify droplets as positive or negative for each given target, select the colored box that best represents the location of the cluster of droplets.
**Direct Quantification with the Advanced Classification Method**

The Advanced Classification Method analysis mode allows the user complete control of droplet identification, and does not force or preassign any clusters. This allows a higher degree of multiplexing through radial dye mixing.

**Important:** This mode is intended for experienced users of droplet digital PCR.

**Setting Up Experiment Parameters**

**To use the advanced classification method**

1. Open the analysis file and select the Plate Editor tab.

2. Select one or more wells to be assigned well analysis attributes.

3. Set the experiment type to Direct Quantification (DQ) and tap or click Apply.

4. (Optional) Change the sample description identifiers.

5. (Optional) Change the sample type. The default type is Unknown.

   **Note:** You cannot change the supermix that was used during data collection.

6. Tap or click Apply.

7. Under Assay Type, select Advanced Classification Method and tap or click Apply.

8. Use the + icon to add up to ten targets.

9. Enter the target name and type (unknown or reference) for each target in the well.
10. (Optional) If you select Reference, you can change the number of reference copies. The default is 2.
   - If no reference is selected, the target in the highest channel is used as the reference.
   - If more than one reference is selected in the well, checkboxes appear to the right for using the reference in the plot.

   **Note:** Each Reference target enables ratio calculations in the results.

11. Assign the first target an expected signal in a channel.

12. Assign your last target a signal in a channel that was not selected for your first target.
   Each target in between will be assigned a signal depending on mixture ratio of the target.

13. Tap or click Apply to confirm all well settings.

   **Tip:** Optionally, to expedite plate setup you can copy and paste data from one well to another, and you can tap or click Undo to undo an applied well setting.

**Viewing and Adjusting Results**

1. Tap or click the 2D Amplitude tab.

   All droplets are unclassified by default and are colored red. Only the Threshold Cluster mode is available for setting thresholds in Advanced Classification experiments in the 2D plot.

2. Use one of the Threshold Cluster mode tools (box, circle or freehand) to assign droplet clusters to targets.

   When you select a droplet cluster, a cluster assignment box appears.

3. Select a radio button to identify the composition of the cluster.

   For example, if the cluster is composed of positive droplets for Target-2, select the positive radio button for Target-2 and the negative radio buttons for the remainder.

   **Important:** For calculations to occur, you must have at least one positive cluster for each target, as well as the assigned negative cluster.
Note: The software sets aside unassigned for calculation purposes. If you are unsure of droplet/cluster composition, leave it unassigned (red).

You can view the concentration results in the data table and in the Concentration tab.

If one of your samples was identified as a Reference under AssayType, you can view a ratio of concentration for unknown versus reference in the Ratio tab.
Appendix B Managing Users

Use the User Setup and Preferences window to add, change, or remove QX Developer users, and set or change user privileges and preferences.

A superuser account, with all permissions, should be set up first. You will use the superuser account to create additional users and assign user privileges. If you need assistance, contact Bio-Rad Technical Support.

**Note**: Users who are not assigned any of the specified privileges can still execute runs and use the Analysis module.

You can set up local or domain accounts. If the computer connected to the QX200 Droplet Reader

- Can access your company’s network, QX Developer can recognize the Active Directory user names.
- Cannot access your company’s network, your IT department can set up user names as local Windows accounts.

You can create user accounts with any user name convention and password character structure, as long as it complies with your organizational requirements. The software is designed to validate each user against the name it finds in either the Active Directory or local domain, so users must log in with the proper domain and user name.
Appendix B Managing Users

However, QX Developer must be able to validate the user in either the local or domain user group before you can add the user in the software. Note the following:

- For local IDs, you must set up each user as a local Windows user account on each computer where the software is installed.
- For Active Directory IDs, your IT department must connect the computer connected to the QX200 Droplet Reader, as well as any separate computers running the software, to your network.

Adding Users

Only a user assigned the Add/Manage Users privilege can add QX Developer users.

Before you begin, ensure that all users have IDs set up in Active Directory or as local accounts.

To add a user

1. In the User Name field, enter the user name and then tap or click Check Name.

   One of the following occurs:

   - If the software recognizes the user name, the software displays a validation message. Continue to Step 2.

   - If the user is not recognized, an error message appears. Do one of the following:

     - For network users, ensure there is a working connection to the network and then verify your entry is accurate.

     - For local users, verify the entry is accurate. If the user is not set up locally or on the network, contact your system administrator for assistance.

2. When the successful validation message appears, tap or click OK, and then tap or click Add.

   The software displays a confirmation message.

3. Tap or click Yes to add the user, and then tap or click OK to close the dialog box.
## Adding or Removing User Privileges

Only a user assigned the Add/Manage users privilege can add or remove user privileges.

By default, individual users without the privileges described in Table 37 can set up and execute a run, open files, view and analyze results.

### Table 37. User Privileges

<table>
<thead>
<tr>
<th>Privilege</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Add/manage users</td>
<td>Add or remove users, set privileges, and change preferences.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Only the superuser can remove this privilege from other users.</td>
</tr>
<tr>
<td>Create new templates</td>
<td>Save a plate, thermal cycling protocol, or analysis report design as a template.</td>
</tr>
<tr>
<td>View data files</td>
<td>View files created by other users in the storage folders.</td>
</tr>
<tr>
<td>created by other users)</td>
<td></td>
</tr>
<tr>
<td>Overwrite existing data file name</td>
<td>Use Save or Save As capability.</td>
</tr>
<tr>
<td></td>
<td>■ Selecting Save replaces the original file content with any changes made by the user, without changing the file name.</td>
</tr>
<tr>
<td></td>
<td>■ Selecting Save As saves the file with a new file name.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Users without this permission can open files and perform analysis but cannot save their changes.</td>
</tr>
<tr>
<td>System settings</td>
<td>View event logs, and view and modify the shared data file and template locations.</td>
</tr>
<tr>
<td></td>
<td>Two locations can be assigned for data files and one for templates.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> You can also set preferred locations, which override all paths in individual preferences.</td>
</tr>
<tr>
<td>Maintenance</td>
<td>Perform software updates.</td>
</tr>
<tr>
<td></td>
<td>View the event log.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> All users can view the maintenance log and maintenance reports.</td>
</tr>
<tr>
<td>Data archive</td>
<td>Move raw data from the computer connected to the QX200 Droplet Reader to free up disk space needed for runs on the instrument.</td>
</tr>
</tbody>
</table>
Table 38 contains a sample set of privileges assigned by user roles common to a laboratory. User roles can be defined and used in any scenario, but may be required in regulatory environments.

Table 38. Example User Roles

<table>
<thead>
<tr>
<th>Privilege</th>
<th>Superuser</th>
<th>Lab Manager</th>
<th>Group Lead</th>
<th>Technician/Student Intern</th>
</tr>
</thead>
<tbody>
<tr>
<td>Add/manage users</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Create new templates</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓*</td>
</tr>
<tr>
<td>View data files created by other users</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Override existing ddPCR file name</td>
<td>✓</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>System settings</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maintenance</td>
<td>✓</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Archive data files</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Rerun a plate for data recovery</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

*You can assign this privilege to non-administrative roles if anyone can create and save a template. If template formats are restricted, then you should assign this permission to administrative roles only.

**Note:** Bio-Rad recommends defining at least one additional superuser as a backup, as well as other administrative and standard roles based on your needs. Guest users can open shared templates, execute runs, and perform analysis but are not typically assigned other privileges.

**To add or modify privileges**

1. Tap or click the Add/Manage Users tab to open the User Management window.
2. In the User Name field, enter a user name.
3. Select or clear the checkbox for each privilege in accordance with the user's role in using the software.

You can assign user privileges in any combination.

4. When the confirmation message appears, tap or click Yes to save the changes, then tap or click OK to close the pop-up.
Modifying User Preferences

Users assigned the Add/Manage Users privilege can change the preferences for other users. Individual users can also change their own preferences.

To modify a user's preferences

1. Tap or click the Add/Manage Users tab, and then enter the user name.
2. Change any of the following preferences for the user:
   - Enter a different file path for user data files and templates.  
     Important: You can also specify Preferred Locations in System Settings, which override all file paths specified in individual preferences. For information, see System Settings Tab on page 194.
   - Select or clear the checkbox to change user data file and template privacy settings.
   - Enter a different default system timeout period.
   - Enter a different number of completed plates to show in the Status section, up to a maximum of 100.
3. Tap or click Save.
   QX Developer displays a confirmation message.
4. Tap or click Yes to save the changes, and then tap or click OK to close the dialog box.

Removing Users

Only a user assigned the Add/Manage Users privilege can remove users. Removing a user from the software does not remove the user from other systems or databases. The user still exists on your network, or in the Windows local user list.

To remove a user

1. Tap or click the Add/Manage Users tab to open the User Management window.
2. In the Current Users pane, select the user and tap or click Remove.
   QX Developer displays a confirmation message.
3. Tap or click Yes to remove the user, and then tap or click OK to close the dialog box.
Appendix B Managing Users
Appendix C Instrument Maintenance

Bio-Rad recommends regular maintenance of your equipment, which includes

- Keeping the surface areas clean
- Regularly checking the fluid levels
- Inspecting the equipment for damaged parts or wiring

Bio-Rad also recommends semi-annual preventive maintenance on your instrument by a Bio-Rad service engineer. Contact Bio-Rad if your instrument requires service or recalibration.

Fluid Maintenance

You must ensure there is enough oil for at least one full run, and you must empty the waste container when it approaches a full state.

You can check your fluid levels using any of the following methods:

- Check the QX Developer status bar, which shows oil as a percentage remaining and waste as percentage full
- Check the Indicator lights on the QX200 Droplet Reader, which flash amber when oil is low and waste is high
- Check the levels in the containers themselves

Table 39 explains fluid level maintenance for your instrument.
Table 39. Fluid Levels

<table>
<thead>
<tr>
<th>QX200 Indicator Light Color</th>
<th>Fluid Level</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid green</td>
<td>Oil greater than 30% or waste less than 70%</td>
<td>Bottle levels OK. You can start your run.</td>
</tr>
<tr>
<td>Flashing green</td>
<td>Oil less than 30% or waste greater than 70%</td>
<td>You may be able to start a run on fewer than 96 wells. If there is not enough oil in the dispensers to perform a run on the number of plates you set up in the Plate Editor, the software prevents the run until you fill the oil dispensers and empty the waste container.</td>
</tr>
<tr>
<td>Flashing amber</td>
<td>Oil less than 10% or waste greater than 90%</td>
<td>The software prevents the run until you fill the oil dispenser.</td>
</tr>
</tbody>
</table>

Replacing Droplet Oil

To replace the oil bottle

1. Slide the container carriage out using the handle on the side of the instrument.
2. Remove the oil bottle and set it aside, to use when changing the waste bottle.
3. Insert a new bottle of oil, and tighten the cap into place.
Managing Waste

You must apply standard Material Safety Data Sheet (MSDS) and OSHA practices when handling and disposing of generated waste. A typical waste profile should contain the following:

- Fluorinated oils: 95%
- Water: 5%
- Bleach: Less than 0.5%
- Proteins, nucleic acids, fluorescent dye: Less than 0.1%

Before disposal, Bio-Rad recommends collecting the waste bottles in a polyethylene container and discarding the container within one month of the oldest waste collected.

While Bio-Rad droplet reader fluids are nonflammable and inert, with low toxicity and environmental impact, they are based on fluorinated hydrocarbon chemistry and disposal should comply with institutional, state, and local regulations. Contact your local environmental health and safety specialists for guidance on proper disposal, including locations in your area.

Use the last empty oil bottle as the new waste bottle. Bio-Rad provides a new Waste label to cover the Oil label.

To replace the waste bottle

1. Slide the container carriage out using the handle on the side of the instrument.
2. Remove the waste bottle.
3. Place the new Waste label over the Oil label on the oil bottle.
4. Insert the last empty oil bottle into the waste container slot.

**Important:** Use each empty oil bottle as a new waste bottle only once and then discard it. Before discarding, add 50ml of 10% bleach solution to the bottle to prevent microbial growth, and then label the bottle as waste.
Flushing or Priming the Instrument

You should flush the instrument after you replace the waste bottle.

**To flush the instrument**

1. Select the System Utilities tab.
2. Tap Flush.
   A message appears, asking if you want to flush the instrument.
3. Tap Yes to begin.
4. A “Performing Flush” message appears.
   When the flushing process is finished, an operation complete message appears.
5. Tap OK.

**To prime the instrument**

You should prime the instrument after you replace the oil bottle.

1. Select the System Utilities tab.
2. Tap Prime.
   A message appears, asking if you want to prime the instrument.
3. Tap Yes to begin.
4. A “Performing Prime” message appears.
   When the priming process is finished, an operation complete message appears.
5. Tap OK.
## Instrument Maintenance Log

<table>
<thead>
<tr>
<th>Month</th>
<th>Year</th>
<th>Date</th>
<th>Daily</th>
<th>Weekly</th>
<th>Monthly</th>
<th>Annually</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Weekly Tasks
- Check filters and filter system
- Check sample loop and rinse
- Check the valve block
- Check and clean the reference electrode

### Monthly Tasks
- Check the electrical connections
- Check the calibration
- Check the sensor

### Annually Tasks
- Check the instrument
- Check the system
- Check the reference solution
- Check the instrument alignment
- Check the system alignment
- Check the sensor alignment
Appendix D System Utilities

From the System Utilities tab, you can access the following tabs for managing tasks and data related to the instrument, software, data, and storage:

- **System Settings** — View shared template and data file locations
- **Event Log** — View logged data about all software activities
- **Maintenance Log** — Record maintenance activities
- **Maintenance Reports** — PDF maintenance records from Bio-Rad maintenance
- **Archive Data** — Clear up space in your primary storage areas
- **Reprocess Data** — Reprocess a .data file using new calibration values
- **Tools** — Flush or prime the instrument

Notes:

- The Instrument Calibration tab is enabled only for the Bio-Rad service engineer user account.
- If you are not connected to the QX200 Droplet Reader, only the System Settings and Event Log tabs are available.
System Settings Tab

The System Settings window opens by default when you select the System Utilities tab, and contains fields where you can enable and enter storage file paths.

In System Settings, you can set up the following storage areas:

- Preferred Location
- Shared Settings

**Important:** Primary storage locations are either the file paths specified in the user's individual preferences, or the file paths specified under Preferred Locations. If a system administrator enables the Preferred Location settings for all users, files paths in individual user preferences are overridden.

**Preferred Location**

If Preferred Location file paths are used, then data files for all users are automatically saved to the specified folders on the instrument.

**Important:** Only users assigned the System Settings user privilege can set preferred locations. This setting overrides individual user storage settings.

**Note:** Preferred location restrictions apply to data files only; users are able to choose a location for template storage.
To specify preferred locations

1. Select the Use for all users checkbox and tap or click Save.
   The file paths are identified in the Datafile Storage Location and Template Storage Location fields.

2. (Optional) Modify the file paths, and then tap or click Save.

Shared Settings

Under Shared Settings, you can specify two shared data file storage folders and one shared template storage folder.

Important: Only users assigned the System Settings user privilege can change storage locations in the System Settings window.

Tip: Because data files are often very large, the software allows you to set a primary storage folder and a backup storage folder. Bio-Rad recommends routinely checking available storage space and if necessary, using the Archive function.

To specify shared folders

1. In the Datafile Storage Location field, enter a file path or tap or click Browse to search for the folder.

2. In the Datafile Storage Location 2 folder, enter a file path or tap or click Browse to search for the folder.
   
   Note: These paths point to secondary storage locations. Data files are always saved initially to the preferred location (if specified by your administrator), or to the path specified in the user’s preferences.

3. In the Template Storage Location field, enter a file path or tap or click Browse to search for the folder.

4. Tap or click Save.
Appendix D System Utilities

System Log Files

From the System Utilities tab, you can access log files for system and software events (event log), and maintenance activities (maintenance log).

From the regulatory edition, you can also generate the audit log as a PDF report.

**Note:** The Audit Log checkbox appears in the list of reportable items in the standard edition, but is disabled.

Event Log

Users who are assigned the Maintenance user privilege can view the system event log, which contains timestamped information on all software activities.

To view the Event Log

1. Select the System Settings tab.

2. Select Event Log and do one of the following:
   - Scroll through the list of events displayed in the Event Log window.
   - Tap or click Open Log and open the text file in Notepad.

3. (Optional) To automatically generate and display a PDF document, tap or click PDF Report.
**Tip:** To restrict access to the file, you can assign a password. Select the PDF Password check box and then enter a password in the field. The file is saved in the Log Reports folder and can be opened only by entering the password.

**Maintenance Log**

All users can view the maintenance log, which contains a list of maintenance records.

Users who are assigned the Maintenance user privilege can set up maintenance records, specify associated activities, and generate PDF reports.

To view the Maintenance Log

1. Select the System Settings tab.

2. Select Maintenance Log.

3. To view the log entries, scroll through the list in the grid.

To add a new activity

1. Tap or click the plus sign (+) next to New Activity.
A line item appears in the grid and the Date Time, User, Product Version, and Firmware Version fields are automatically completed.

2. In the Activity field, enter a description of the planned activity.
3. Tap or click Save.
4. Repeat to add another activity.

After tapping or clicking Save, you can create a PDF report of the added activities.
5. (Optional) Tap or click PDF Report.

**Maintenance Reports**

Reports generated from the Bio-Rad service engineer when performing maintenance on your instrument are stored on the Maintenance Reports tab.

► To open a report, tap or click the list item.

Following is an example of a Color Calibration report:
Archiving Data

When the storage space on the computer connected to the QX200 Droplet Reader approaches a predefined limit, the software displays an advisory message at set intervals. Users can free up space by moving raw data to an archive storage folder in a different location.

**Important:** You must be assigned the Data Archive user privilege to archive data.

**To archive files**

1. Select the System Settings tab.

2. Select Archive Data.

3. Select a tab to archive a file type. For example, select Raw Data.

4. Under Raw Data Archive Progress, tap or click the calendar icons and select a From date and a To date to define a date span.

5. Click Query to search for files dated within the date span.

6. Tap or click Browse to search for a select the target storage location.

7. Tap or click Archive.
Reprocessing Data

After your instrument has been recalibrated, you can reprocess a data file from a run using different color calibration values.

**Note:** The instrument must be in an idle state before you can reprocess the data in QX Developer. If the file was created in an older software version, the software cannot reprocess the file and displays an advisory message.

**To reprocess a data file**

1. Select the System Utilities tab.

2. Tap Reprocess Data.

3. Tap Browse and navigate to the analysis data file, and then in the Select Datafile dialog box, select the file and tap or click Open.

   The calibration values appear in the first Color Calibration table.

4. Under the New calibration values heading, enter new values for the FAM, HEX, VIC, and EvaGreen® fluorophores.

5. Tap Reprocess Data.

   **Note:** When you reprocess a data file, the software creates an audit record containing the old and new calibration values.
System Calibration and Tools

You cannot execute runs on the QX200 Droplet Reader unless the instrument is properly calibrated.

Only a Bio-Rad service engineer can calibrate your instrument and use the Instrument Calibration options in QX Developer.

If your instrument requires service, contact Bio-Rad Technical Support.

The Tools functions are currently applicable only to instrument maintenance. For information, see Flushing or Priming the Instrument on page 190.
Appendix D System Utilities
Appendix E Additional Computers

For analysis purposes only, you can download purchased software from the Bio-Rad website and install it on computers provided by your company. This section explains the recommended system requirements for QX Developer, and provides instructions for downloading and installing or updating the software.
Appendix F Troubleshooting

This section contains information on troubleshooting and correcting minor issues you may experience with your instrument or software.

Event Log

The Event Log contains records for all system and user activities, including system errors. If you do not understand, or cannot find an immediate remedy to, the error event, contact Bio-Rad Technical Support.

Instrument Errors

If an error occurs during a run, the software cancels the run and displays an X in the Status column, along with a Run Aborted by System message as the Run End status. Appendix F contains error grouped by type.

1. Tap the X to display the corresponding error code and description.
2. Perform the steps in the solution column before starting a new run.
3. If the suggested solution does not resolve the problem, contact Bio-Rad Technical Support immediately.

   **Note:** The error code and description are also displayed in the Run Information window.

Communication errors can be caused by instrument, connectivity, or power failures.
Appendix F Troubleshooting

Table 40. Communication errors

<table>
<thead>
<tr>
<th>Error Code</th>
<th>Description</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>102, 103, 104*, 110*, 111</td>
<td>Communication loss between the instrument and software, detector board error, loss during a run, or unable to reestablish a connection</td>
<td>Disconnect and reconnect the USB cable, and then power cycle the instrument.</td>
</tr>
<tr>
<td>Note: For codes marked with an asterisk (*)</td>
<td>some data may be usable.</td>
<td></td>
</tr>
<tr>
<td>302, 308</td>
<td>Communication error involving pump</td>
<td>Power cycle the instrument.</td>
</tr>
<tr>
<td>Note: Some data might be usable.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>502</td>
<td>Invalid checksum value</td>
<td></td>
</tr>
<tr>
<td>900</td>
<td>Invalid command</td>
<td></td>
</tr>
</tbody>
</table>

An electrical error is typically triggered by an invalid setting.

Table 41. Electrical error codes

<table>
<thead>
<tr>
<th>Error Code</th>
<th>Description</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>406</td>
<td>Electrical Error; invalid power value setting.</td>
<td>Power cycle the instrument.</td>
</tr>
<tr>
<td>Note: Some data might be usable.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Mechanical errors can be caused by fluid issues (low oil, high waste), instrument operation errors, or binary data collection errors.

Table 42. Mechanical error codes

<table>
<thead>
<tr>
<th>Error Code</th>
<th>Description</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>105, 106</td>
<td>Fluidics or motion response was invalid.</td>
<td>Power cycle the instrument.</td>
</tr>
<tr>
<td>Note: Some data may be usable.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>201</td>
<td>Motors disabled</td>
<td>Remove and reinsert the plate and plate lid.</td>
</tr>
</tbody>
</table>
1. |                                                                                                           |
2. | Check for incorrect or multiple heat seal foil layers.                                               |
3. | Check for strain on all motor axes.                                                                |
4. | Power cycle the instrument.                                                                      |
Table 42. Mechanical error codes, continued

<table>
<thead>
<tr>
<th>Error Code</th>
<th>Description</th>
<th>Solution</th>
</tr>
</thead>
</table>
| 205, 207, 210, 211, 212, 218 | X, Y, or Z-axis, unable to reach target location or home in allotted time.  
**Note:** Some data might be usable. | Check for motor obstructions, and then power cycle the instrument. |
| 301, 303, 304, 305, 307, 312, 316, 317 | Pump errors  
**Note:** Some data might be usable. | Power cycle the instrument. |
Appendix G Ordering Information

This appendix contains descriptions and catalog numbers for new or replacement instruments, accessories, and consumables for Bio-Rad's ddPCR products, including your QX200 Droplet Reader.

ddPCR System and Instrument Packages

Table 43 contains information on the full QX200 Droplet Digital PCR system, as well as individual ddPCR instruments and included accessories.

Table 43. ddPCR System and Instrument Packages

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>QX200 Droplet Digital PCR System</td>
<td>Includes the QX200 Droplet Generator, QX200 Droplet Reader, computer, software, and consumables</td>
<td>1864001</td>
</tr>
<tr>
<td>QX200 Droplet Generator</td>
<td>Includes the droplet generator (1), cartridges and gaskets (24 each), cartridge holders (2), and power cord (1)</td>
<td>1864002</td>
</tr>
<tr>
<td>QX200 Droplet Reader</td>
<td>Includes the droplet reader (1), plate holders (2), USB cable (1), and power cord (1)</td>
<td>1864003</td>
</tr>
<tr>
<td>Automated Droplet Generator</td>
<td>Includes the automated droplet generator (1), cooling block accessory (1), oil purge reservoir (1), and power cord (1)</td>
<td>18641001</td>
</tr>
<tr>
<td>C1000 Thermal Cycler</td>
<td>Touch thermal cycler with 96-deep well reaction module</td>
<td>1851197</td>
</tr>
</tbody>
</table>
### QX200 Droplet Reader Accessories

Table 44 contains information on accessories for the QX200 Droplet Reader.

**Table 44. QX200 Droplet Reader Accessories**

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Computer and QX Developer software</td>
<td>Computer that connects to the QX200 Droplet Reader for data collection and analysis</td>
<td>17006483</td>
</tr>
<tr>
<td>USB cable and power cord</td>
<td>Cable connecting the computer to the instrument and power cord connecting the instrument to the power source</td>
<td>Included</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Contact Bio-Rad Technical Support for replacement information.</td>
<td></td>
</tr>
<tr>
<td>Droplet reader plate holders (2)</td>
<td>Used to position the 96-well plate in the droplet reader plate compartment</td>
<td>12006834</td>
</tr>
</tbody>
</table>

**Note:** Reagents include the PCR supermix (for probes or EvaGreen®), droplet generation oil, and buffer control kit (for probes or EvaGreen®).

### QX200 Droplet Generator Accessories

Table 45 contains information on the accessories for the QX200 Droplet Generator.

**Table 45. QX200 Droplet Generator Accessories**

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>DG8 droplet generator cartridges and gaskets</td>
<td>Microfluidic cartridges used to mix sample and oil for droplet generation</td>
<td>1864007</td>
</tr>
<tr>
<td></td>
<td>Gaskets to seal the cartridges</td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> You must seal the cartridge to prevent evaporation and to apply required pressure for droplet generation.</td>
<td></td>
</tr>
<tr>
<td>Droplet generator cartridge holder</td>
<td>Positions and holds the droplet generator cartridge in the instrument compartment</td>
<td>1863051</td>
</tr>
<tr>
<td>Power cord</td>
<td>Connects the QX200 Droplet Generator to the power source</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> Contact Bio-Rad Technical Support for replacement information.</td>
<td></td>
</tr>
</tbody>
</table>
Automated Droplet Generator Accessories

Table 46 contains information on accessories for the Automatic Droplet Generator.

Table 46. Automated Droplet Generator Accessories

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cooling block accessory</td>
<td>Prevents evaporation during droplet generation</td>
<td>12002819</td>
</tr>
<tr>
<td>DG32 droplet generator cartridges</td>
<td>Microfluidic cartridge with gasket for mixing sample and droplet generation oil</td>
<td>1864108</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1864109</td>
</tr>
<tr>
<td>Oil purge reservoir</td>
<td>Collects oil waste from priming and flushing</td>
<td></td>
</tr>
<tr>
<td>Power cord</td>
<td>Connects the Automated Droplet Generator to the power source</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Note: Contact Bio-Rad Technical Support for replacement information.

ddPCR Materials and Consumables

Table 47 contains information on accessories and consumables for Bio-Rad instruments.

Table 47. Consumables and Other Materials

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>96-well PCR plates</td>
<td>ddPCR 96-well plates</td>
<td>12001925</td>
</tr>
<tr>
<td>DG32 droplet generator cartridges (AUTODG)</td>
<td>Microfluidic cartridge with gasket for mixing sample and droplet generation oil in the Automated Droplet Generator</td>
<td>1864108</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1864109</td>
</tr>
<tr>
<td>Rainin Pipets</td>
<td>20 µL for sample loading</td>
<td>L-20, L8-20</td>
</tr>
<tr>
<td></td>
<td>50 µL for droplet transfer</td>
<td>L-50, L8-50</td>
</tr>
<tr>
<td></td>
<td>8-channel, 200 µL for oil</td>
<td>L8-200</td>
</tr>
<tr>
<td>Rainin Pipet tips</td>
<td>Filtered</td>
<td>GP-L10F</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GP-L200F</td>
</tr>
<tr>
<td>Pipet tips (AUTODG)</td>
<td>Tips for Automated Droplet Generator</td>
<td>1864120</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1864121</td>
</tr>
</tbody>
</table>
### Table 47. Consumables and Other Materials, continued

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pipet tip bin (AUTODG)</td>
<td>Tip waste bins for Automated Droplet Generator</td>
<td>1864125</td>
</tr>
<tr>
<td>Foil plate seals</td>
<td>Pierceable foil plate seals</td>
<td>1814040</td>
</tr>
<tr>
<td>Plate sealer</td>
<td>PX1 PCR plate sealer</td>
<td>1814000</td>
</tr>
<tr>
<td>Droplet generation oils</td>
<td>Droplet generation oil for probes</td>
<td>1863005</td>
</tr>
<tr>
<td></td>
<td>Automated droplet generator oil for EvaGreen®</td>
<td>1864112</td>
</tr>
<tr>
<td></td>
<td>Automated droplet generator oil for probes</td>
<td>1864110</td>
</tr>
<tr>
<td>Buffer controls</td>
<td>QX200 buffer control kit for EvaGreen®</td>
<td>1864052</td>
</tr>
<tr>
<td></td>
<td>QX200 buffer control kit for probes</td>
<td>1863052</td>
</tr>
<tr>
<td>Droplet reader oil</td>
<td>ddPCR droplet reader oil</td>
<td>1863004</td>
</tr>
<tr>
<td>Droplet reader waste bottle</td>
<td>You can use an empty droplet reader oil bottle to collect the waste from droplet reading.</td>
<td>N/A</td>
</tr>
</tbody>
</table>

**PCR Supermixes**

**Note:** You can use these supermixes in both the QX200 Droplet Generator and the Automatic Droplet Generator.

<table>
<thead>
<tr>
<th>Name</th>
<th>Catalog Number</th>
<th>Volume (ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>QX200 ddPCR EvaGreen® supermix</td>
<td>1864033</td>
<td>2ml (2 x)</td>
</tr>
<tr>
<td>For use in nucleic acid sample preparation with the QX200 droplet generator</td>
<td>1864034</td>
<td>1ml</td>
</tr>
<tr>
<td></td>
<td>1864035</td>
<td>5ml (5 x)</td>
</tr>
<tr>
<td></td>
<td>1864036</td>
<td>1ml</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25ml (25 x)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1ml</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50ml (50 x)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1ml</td>
</tr>
<tr>
<td>ddPCR Supermix for Probes (no dUTP)</td>
<td>1863023</td>
<td>2ml (2 x)</td>
</tr>
<tr>
<td>2x supermix, for use in nucleic acid sample preparation with the QX100 or QX200 droplet generators</td>
<td>1863024</td>
<td>1ml</td>
</tr>
<tr>
<td></td>
<td>1863025</td>
<td>5ml (5 x)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1ml</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25ml (25 x)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1</td>
</tr>
</tbody>
</table>
### Table 47. Consumables and Other Materials, continued

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>ddPCR Supermix for Probes</td>
<td></td>
<td>1863026</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> 2x supermix, for use in sample preparation with the QX100 or QX200 droplet generators</td>
<td>1863010, 5ml (5 x 1ml)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1863027</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1863028</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25ml (25 x 1ml)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50ml (50 x 1ml)</td>
</tr>
<tr>
<td>ddPCR Supermix for Residual DNA Quantification</td>
<td></td>
<td>1864037</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> 2x supermix, for use in residual DNA detection with the QX100 or QX200 droplet generators</td>
<td>1864038</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1864039</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1864040</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25ml (25 x 1ml)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50ml (50 x 1ml)</td>
</tr>
<tr>
<td>One-step RT-ddPCR Advanced Kit for Probes</td>
<td></td>
<td>1864033</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> 200 or 500 x 20 µl reactions, 2x RT0-ddPCR mix, includes 1 or 2 manganese acetate tubes</td>
<td>1864034</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2ml (2 x 1ml)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5ml (5 x 1ml)</td>
</tr>
<tr>
<td><strong>ddPCR Mutation Screening Kits</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ddPCR KRAS G12/G13 Screening Kit</td>
<td></td>
<td>1863506</td>
</tr>
<tr>
<td>ddPCR KRAS Q61 Screening Kit</td>
<td></td>
<td>12001626</td>
</tr>
<tr>
<td>ddPCR BRAF V600 Screening Kit</td>
<td></td>
<td>12001037</td>
</tr>
<tr>
<td>ddPCR NRAS Q61 Screening Kit</td>
<td></td>
<td>12001006</td>
</tr>
<tr>
<td>ddPCR NRAS G12 Screening Kit</td>
<td></td>
<td>12001094</td>
</tr>
<tr>
<td>ddPCR NRAS G12/G13 Screening Kit</td>
<td></td>
<td>12001627</td>
</tr>
<tr>
<td>ddPCR EGFR Exon 19 Deletions Screening Kit</td>
<td></td>
<td>12002392</td>
</tr>
<tr>
<td><strong>Note:</strong> All kits include 20x multiplex assay and 2x ddPCR supermix for probes (no DUTP)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| **ddPCR Residual DNA Quantification Kits**   |                                                               |                |

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Table 47. Consumables and Other Materials, continued

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>ddPCR CHO Residual DNA Quantification Kit</td>
<td></td>
<td>17000031</td>
</tr>
<tr>
<td>ddPCR E. coli Residual DNA Quantification Kit</td>
<td></td>
<td>17000032</td>
</tr>
<tr>
<td><strong>Note:</strong> 200 x 20 µl reactions, includes 20x CHO or E.coli RDQ assay and 2X ddPCXR Supermix for Residual DNA Quantification</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**ddPCR Copy Number Determination Kits**

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>ddPCR SMN1 Copy Number Determination Kit</td>
<td></td>
<td>1863500</td>
</tr>
<tr>
<td>ddPCR SMN2 Copy Number Determination Kit</td>
<td></td>
<td>1863503</td>
</tr>
<tr>
<td><strong>Note:</strong> 200 x 20 µl reactions, includes assay at 20x concentration, 2x ddPCR Supermix for Probes (no dUTP) and positive controls</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**ddPCR Library Quantification Kits**

<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>ddPCR Library Quantification Kit for Illumina TruSeq</td>
<td></td>
<td>1863040</td>
</tr>
<tr>
<td><strong>Note:</strong> 200 x 20 µl reactions, includes 1 vial of primers and probes at 20x concentration, 2x ddPCR Supermix for Probes (no dUTP) and positive controls</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**ddPCR Genome Edit Detection Assays**
<table>
<thead>
<tr>
<th>Product</th>
<th>Description</th>
<th>Catalog Number</th>
<th>rxns</th>
</tr>
</thead>
<tbody>
<tr>
<td>ddPCR HDR Gene Edit Assay</td>
<td></td>
<td>12002312</td>
<td>100</td>
</tr>
<tr>
<td>ddPCR HDR Gene Edit Assay</td>
<td></td>
<td>12002313</td>
<td>rxns</td>
</tr>
<tr>
<td>ddPCR HDR Gene Edit Package</td>
<td></td>
<td>12003796</td>
<td>500</td>
</tr>
<tr>
<td>ddPCR HDR Ref Assay, Predesigned</td>
<td></td>
<td>12003805</td>
<td>rxns</td>
</tr>
<tr>
<td>ddPCR HDR Ref Assay, Predesigned</td>
<td></td>
<td>12003806</td>
<td>1,000</td>
</tr>
<tr>
<td>ddPCR HDR Ref Package, Predesigned</td>
<td></td>
<td>12003793</td>
<td>rxns</td>
</tr>
<tr>
<td>ddPCR NHEJ Gene Edit Assay</td>
<td></td>
<td>12002314</td>
<td>100</td>
</tr>
<tr>
<td>ddPCR NHEJ Gene Edit Assay</td>
<td></td>
<td>12002315</td>
<td>rxns</td>
</tr>
<tr>
<td>ddPCR NHEJ Gene Edit Package</td>
<td></td>
<td>12003794</td>
<td>500</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 47. Consumables and Other Materials, continued