

resDNASEQ™ Quantitative DNA Kits

USER GUIDE

for use with:

(Genomic DNA)

- resDNASEQ™ Quantitative CHO DNA Kit
- resDNASEQ™ Quantitative *E. coli* DNA Kit
- resDNASEQ™ Quantitative HEK293 DNA Kit
- resDNASEQ™ Quantitative Human DNA Kit
- resDNASEQ™ Quantitative MDCK DNA Kit
- resDNASEQ™ Quantitative NS0 DNA Kit
- resDNASEQ™ Quantitative *Pichia* DNA Kit
- resDNASEQ™ Quantitative Sf9 and Baculovirus DNA Kit
- resDNASEQ™ Quantitative Vero DNA Kit

(Plasmid DNA)

- resDNASEQ™ Quantitative Plasmid DNA - Kanamycin Resistance Gene Kit

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For descriptions of symbols on product labels or product documents, go to [thermofisher.com/symbols-definition](https://www.thermofisher.com/symbols-definition).

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| Revision | Date | Description |
|----------|---------------|--|
| J | 28 April 2021 | Update to the control serial dilutions required for the resDNASEQ™ Quantitative Plasmid DNA - Kanamycin Resistance Gene Kit (Cat. No. A50337). |
| H | 2 April 2021 | Update to include the resDNASEQ™ Quantitative Plasmid DNA - Kanamycin Resistance Gene Kit (Cat. No. A50337). |

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Product information

IMPORTANT! Before using this product, read and understand the information in the “Safety” appendix in this document.

Product description

The resDNASEQ™ Quantitative DNA Kits are used to quantitate residual DNA from CHO, *E. coli*, HEK293, Human, MDCK, NS0, *Pichia*, Sf9 and Baculovirus, and Vero, or plasmid DNA for Kanamycin resistance, in cell lines which are used for production of biopharmaceutical products. Use the kit after you extract host-cell DNA from test samples. For extraction information, see the *PrepSEQ™ Sample Preparation Kits User Guide* (Pub. No. 4469838).

The resDNASEQ™ Quantitative DNA Kits use TaqMan™ quantitative PCR to perform rapid, specific quantitation of sub-picogram levels of residual host-cell or plasmid DNA. The assay is accurate and reliable across a broad range of sample types, from in-process samples to final product.

To generate the standard curve used to quantitate the DNA in test samples, the CHO, MDCK, NS0, Vero, and Plasmid DNA - Kanamycin resistance assays require six dilutions (from 30 fg to 3 ng per reaction) and the *E. coli*, HEK293, Human, *Pichia*, and Sf9 and Baculovirus assays require five dilutions (300 fg to 3 ng per reaction). Control DNA for standard curve generation is included in the kits. In addition, the kits use an internal positive control (IPC) to evaluate the performance of each PCR reaction.

Contents

Table 1 resDNASEQ™ Quantitative CHO DNA Kit (Cat. No. 4402085)

| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ CHO DNA Control | | |
| CHO DNA Control, 30 ng/μL | 40 μL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |
| resDNASEQ™ CHO Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| 10X CHO DNA Real-Time PCR Assay Mix | 300 μL | -25°C to -15°C, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |

Table 2 resDNASEQ™ Quantitative *E. coli* DNA Kit (Cat. No. 4458435)

| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |
| 10X <i>E. coli</i> DNA Assay Mix | 300 μL | -25°C to -15°C, protect from light |
| resDNASEQ™ <i>E. coli</i> DNA Control | | |
| <i>E. coli</i> DNA Control, 30 ng/μL | 40 μL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |

Table 3 resDNASEQ™ Quantitative HEK293 DNA Kit (Cat. No. A46014)

| Contents | Amount | Storage |
|---|-------------|--|
| resDNASEQ™ HEK293 DNA Control | | |
| HEK293 DNA Control, 30 ng/μL | 40 μL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |
| resDNASEQ™ HEK293 DNA Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| 10X HEK293 Assay Mix | 300 μL | -25°C to -15°C, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |

Table 4 resDNASEQ™ Quantitative Human DNA Kit (Cat. No. A26366)

| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ Human DNA Control | | |
| Human DNA Control, 30 ng/μL | 40 μL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |
| resDNASEQ™ Human Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| 10X Human DNA Assay Mix | 300 μL | -25°C to -15°C, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |

Table 5 resDNASEQ™ Quantitative MDCK DNA Kit (Cat. No. 4464335)

| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |
| 10X MDCK DNA Assay Mix | 300 µL | -25°C to -15°C, protect from light |
| resDNASEQ™ MDCK DNA Control | | |
| MDCK DNA Control, 30 ng/µL | 40 µL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |

Table 6 resDNASEQ™ Quantitative NS0 DNA Kit (Cat. No. 4458441)

| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |
| 10X NS0 DNA Assay Mix | 300 µL | -25°C to -15°C, protect from light |
| resDNASEQ™ NS0 DNA Control | | |
| NS0 DNA Control, 30 ng/µL | 40 µL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |

Table 7 resDNASEQ™ Quantitative *Pichia* DNA Kit (Cat. No. 4464336)

| Contents | Amount | Storage |
|---|-------------|--|
| resDNASEQ™ Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |
| 10X <i>Pichia</i> DNA Assay Mix | 300 µL | -25°C to -15°C, protect from light |
| resDNASEQ™ <i>Pichia</i> DNA Control | | |
| <i>Pichia</i> DNA Control, 30 ng/µL | 40 µL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |

Table 8 resDNASEQ™ Quantitative Sf9 and Baculovirus DNA Kit (Cat. No. A46066)

| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |
| 10X Sf9 + Baculovirus DNA Assay Mix | 300 µL | -25°C to -15°C, protect from light |
| resDNASEQ™ Sf9 and Baculovirus DNA Control | | |
| Multiplex DNA Control with: <ul style="list-style-type: none"> • Sf9 DNA Control, 30 ng/µL • Baculovirus DNA Control, 30 ng/µL | 40 µL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |

Table 9 resDNASEQ™ Quantitative Vero DNA Kit (Cat. No. A41797)

| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ Real-Time PCR Reagents | | |
| 2X Environmental Master Mix | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |
| 10X Vero DNA Assay Mix | 300 µL | -25°C to -15°C, protect from light |
| resDNASEQ™ Vero DNA Control | | |
| Vero DNA Control, 30 ng/µL | 40 µL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |

Table 10 resDNASEQ™ Quantitative Plasmid DNA - Kanamycin Resistance Gene Kit (Cat. No. A50337)

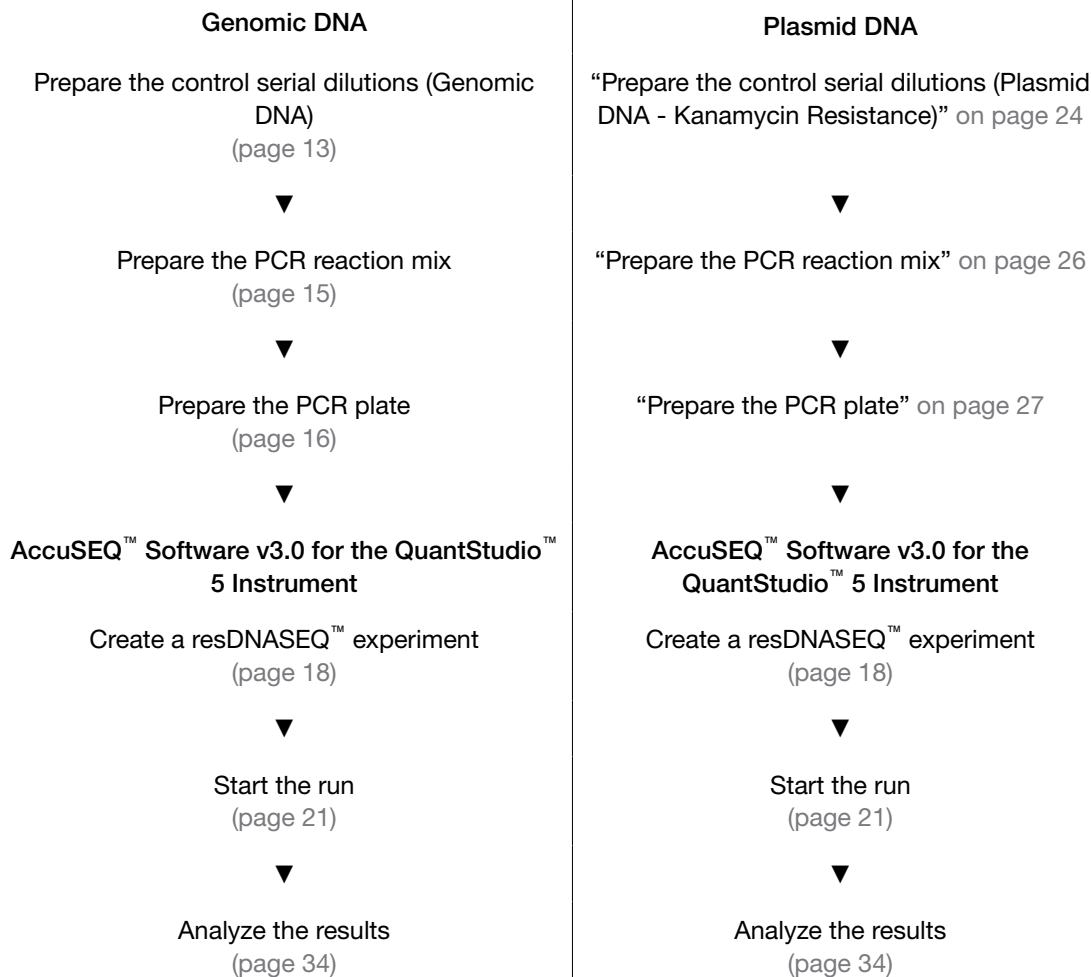
| Contents | Amount | Storage |
|--|-------------|--|
| resDNASEQ™ Kanamycin DNA Control | | |
| KanR DNA Conc: 3 × 10 ⁷ copies/µL | 44 µL | -25°C to -15°C |
| DNA Dilution Buffer (DDB) | 7 mL | 2-8°C |
| resDNASEQ™ Kanamycin DNA Real-Time PCR Reagents | | |
| 2X Environmental Master Mix 3.0 | 2 × 0.75 mL | -25°C to -15°C before first use, protect from light 2-8°C after first use, protect from light |
| 10X KanR Assay Mix | 300 µL | -25°C to -15°C, protect from light |
| Yeast tRNA (10mg/mL) | 500 µL | -25°C to -15°C |
| Negative Control (water) | 1.0 mL | -25°C to -15°C before first use 2-8°C after first use |

Required materials not supplied

Unless otherwise indicated, all materials are available through thermofisher.com. "MLS" indicates that the material is available from fisherscientific.com or another major laboratory supplier.

| Item | Source |
|---|---|
| Instrument | |
| QuantStudio™ 5 Real-Time PCR System with AccuSEQ™ Real-Time PCR Software v3.1 or later | Contact your local sales representative. |
| Consumables | |
| MicroAmp™ Optical 96-Well Reaction Plate with Barcode, 20 plates, 0.1-mL wells; for use with Applied Biosystems™ 7500 Fast Real-Time PCR System | 4346906 |
| MicroAmp™ 96-Well Base | N8010531 , 10 bases |
| MicroAmp™ Optical Adhesive Film | 4311971 , 100 covers 4360954 , 25 covers |
| MicroAmp™ Adhesive Film Applicator | 4333183 , 5 applicators |
| Miscellaneous items | |
| Disposable gloves | Major lab supplier (MLS) |
| Pipettes | MLS |
| Aerosol-resistant micropipette tips | MLS |
| For the PCR plate: Fisher Scientific™ Mini Plate Spinner Centrifuge, 120- or 230-volt | 14-100-143 (120-volt), 14-100-141 (230-volt) |
| Nonstick, RNase-free Microfuge Tubes, 1.5-mL (1 box; 250 tubes/box) | AM12450 |

Workflow





Genomic DNA quantitation

Prepare the control DNA serial dilutions for the standard curve

Guidelines for standard dilutions

- Prepare the standard curve and the test samples in different areas of the lab.
- Use different sets of pipettors for test sample preparation and for standard curve preparation and aliquoting to avoid cross-contamination of test samples.
- Vortex each tube to mix the contents thoroughly before each dilution step.
- Briefly centrifuge to collect all the liquid at the bottom before making the next dilution.

Prepare the control serial dilutions (Genomic DNA)

1. Label nonstick 1.5-mL microfuge tubes: **NTC, SD1, SD2, SD3, SD4, SD5**.
For CHO, Vero, MDCK, and NS0 kits, label an additional tube with **SD6**.
2. Add 50 μL of DNA Dilution Buffer (DDB) to tube NTC. Put aside.
3. Add 990 μL of DNA Dilution Buffer (DDB) to tube SD1.
4. Add 450 μL of DNA Dilution Buffer (DDB) to tubes SD2, SD3, SD4, SD5, and (for CHO, Vero, MDCK, and NS0 only) SD6.
5. Remove the tube of DNA control (30 ng/ μL) from the freezer.
6. After the DNA thaws, vortex it gently for 2 seconds, then briefly centrifuge.
7. Perform the serial dilutions:
 - a. Add 10 μL of the DNA control to the tube that is labeled SD1, then vortex thoroughly and briefly centrifuge.
 - b. Transfer 50 μL of the DNA from tube SD1 to tube SD2, then vortex thoroughly and briefly centrifuge.

- c. Continue to transfer 50 μL of DNA from the previous dilution tube to the next dilution tube until you add DNA to tube SD5 (*E. coli*, HEK293, Human, *Pichia*, and Sf9 and Baculovirus) or SD6 (CHO, Vero, MDCK, and NS0). Final dilutions are shown in the table. After each transfer, vortex thoroughly, then centrifuge briefly.

| Serial dilution (SD) tube | Dilution | pg DNA/reaction (10 μL of the diluted DNA used in final 30 μL of PCR reaction) |
|--|--|--|
| Control | DNA control tube | 300,000 pg |
| SD 1 | 10 μL DNA control + 990 μL DDB | 3,000 pg |
| SD 2 | 50 μL SD1 + 450 μL DDB | 300 pg |
| SD 3 | 50 μL SD2 + 450 μL DDB | 30 pg |
| SD 4 | 50 μL SD3 + 450 μL DDB | 3 pg |
| SD 5 | 50 μL SD4 + 450 μL DDB | 0.3 pg |
| SD 6 (for CHO, Vero, MDCK, and NS0 only) | 50 μL SD5 + 450 μL DDB | 0.03 pg |

8. Store the DNA dilution tubes:

| Temperature | For use |
|-------------|--|
| 4°C | Same day |
| -20°C | ≤ 1 week |
| -20°C | SD1 in single-use aliquots ≤ 6 months |

Prepare the PCR reaction mix

1. Determine the number of reactions needed for the controls and test samples that you will quantify.
2. Thaw all kit reagents completely at room temperature, thoroughly mix reagents, and briefly centrifuge.
3. Prepare a PCR reaction mix using the reagents and volumes shown in the table below.
 - Multiply the PCR reaction volume for one reaction (30 μ L) by the number of reactions that you need to run.
 - Use 10% excess volume to compensate for pipetting losses.

Note: Use reagents from the same lot for all reactions.

| Kit reagents | Volume for 1 30- μ L reaction | Volume for 36 30- μ L reactions (includes 10% overage) |
|--|-----------------------------------|---|
| Negative Control (water) | 2 μ L | 79.2 μ L |
| 10X DNA assay mix appropriate for the cell line being tested | 3 μ L | 118.8 μ L |
| 2X Environmental Master Mix | 15 μ L | 594 μ L |
| DNA template | 10 μ L | Add DNA template to each well separately, not as part of Master Mix |
| Total | 30 μL | 792 μL |

Prepare the PCR plate

Plate setup differs slightly for each AccuSEQ™ System. See your software user guide for specific instructions. Place samples, NTCs, and standards in different quadrants of the plate.

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | | |
|---|--------|-------------------------------------|-------------------------------------|-------------------------------------|--------|--------|--------|--------|--------|--------|---|---|---|---------------------------------------|
| ① | A | NTC [N E ... x] [U IP ... x] | NTC [N E ... x] [U IP ... x] | NTC [N E ... x] [U IP ... x] | Sample | Sample | Sample | Sample | Sample | Sample | STD 3000 [S E ... x] [U IP ... x] | STD 3000 [S E ... x] [U IP ... x] | STD 3000 [S E ... x] [U IP ... x] | |
| | B | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | STD 300 [S E ... x] [U IP ... x] | STD 300 [S E ... x] [U IP ... x] | STD 300 [S E ... x] [U IP ... x] | |
| | C | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | STD 30 [S E ... x] [U IP ... x] | STD 30 [S E ... x] [U IP ... x] | STD 30 [S E ... x] [U IP ... x] |
| | D | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | STD 3 [S E ... x] [U IP ... x] | STD 3 [S E ... x] [U IP ... x] | STD 3 [S E ... x] [U IP ... x] |
| ② | E | TS 1 [U E ... x] [U IP ... x] | TS 1 [U E ... x] [U IP ... x] | TS 1 [U E ... x] [U IP ... x] | Sample | Sample | Sample | Sample | Sample | Sample | STD 0.3 [S E ... x] [U IP ... x] | STD 0.3 [S E ... x] [U IP ... x] | STD 0.3 [S E ... x] [U IP ... x] | |
| | F | TS 2 [U E ... x] [U IP ... x] | TS 2 [U E ... x] [U IP ... x] | TS 2 [U E ... x] [U IP ... x] | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | |
| | G | TS 3 [U E ... x] [U IP ... x] | TS 3 [U E ... x] [U IP ... x] | TS 3 [U E ... x] [U IP ... x] | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | |
| | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | Sample | | |
| H | | | | | | | | | | | | | | |

Figure 1 Default plate setup in the AccuSEQ™ Real-Time PCR Software v3.1

- ① No template controls
- ② Samples
- ③ Standard curve

1. Add 20 µL PCR reaction mix to each well.
2. Add 10 µL of PCR NTC to the appropriate wells.
3. Add 10 µL each of extracted sample DNA to the appropriate wells.

Note: If you prepared samples with the automated protocol, use a multichannel pipette to transfer the extracted sample.

4. Add 10 μ L of standard dilutions to the appropriate wells.

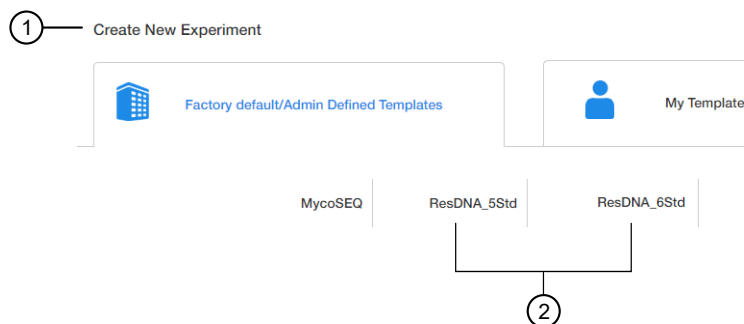
Note: Use different sets of pipettors to dispense test sample and standard curve dilutions to avoid cross-contamination of test samples.

5. Use a film applicator to seal the plate with optical film, then briefly centrifuge with a miniplate centrifuge that is compatible with 96-well plates.

Setup, run, and analyze samples with AccuSEQ™ Software on the QuantStudio™ 5 Real-Time PCR Instrument

Create a resDNASEQ™ experiment

1. In the **Home** screen, click the **Factory default/Admin Defined Template** tab, then select a **resDNASEQ** template.



- ① **Factory default/Admin Defined Template** tab
- ② **resDNASEQ** template (ResDNA_5Std or ResDNA_6Std)

| Serial Dilutions (Standards) | Template | Assays |
|------------------------------|----------|---|
| 5 | _5Std | <i>E. coli</i> , HEK293, Human, <i>Pichia</i> , Sf9 and Baculovirus |
| 6 | _6Std | CHO, Vero, MDCK, NS0 |

2. In the **Experiment Properties** pane of the **Setup** tab:
 - a. (Optional) Change the system-generated name of the experiment.
 - b. (Optional) Enter the plate **Barcode**, then add **Comments**.
Default resDNASEQ™ settings (cannot be changed)
 - **Experiment Type** is **Quantitation-Standard Curve**
 - **Chemistry** is **TaqMan™ Reagents**
 - **Ramp Speed** is **Standard - 2hrs**
 - c. Click **Next**.

Note: Experiment names cannot be changed after this step.

- In the **qPCR Method** pane of the **Setup** tab, view the default volume and cycling conditions (cannot be changed).

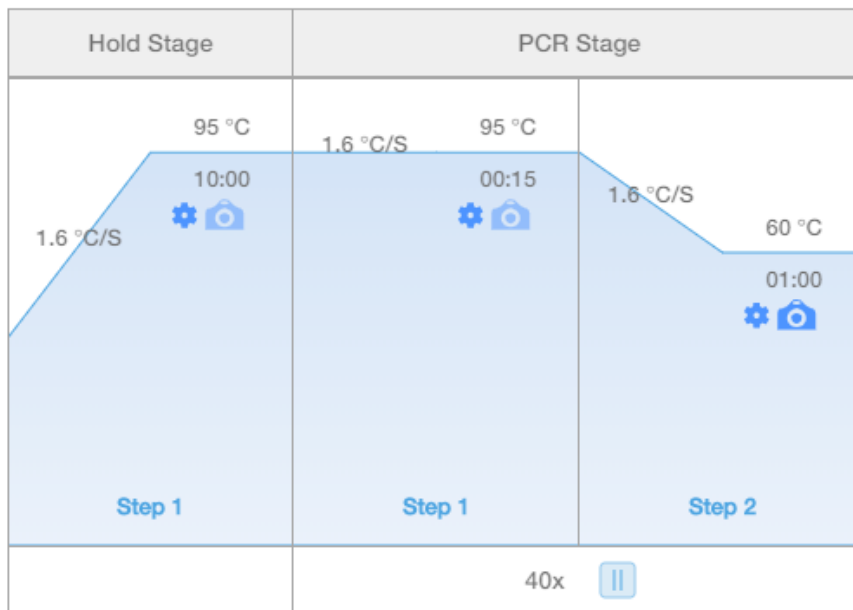
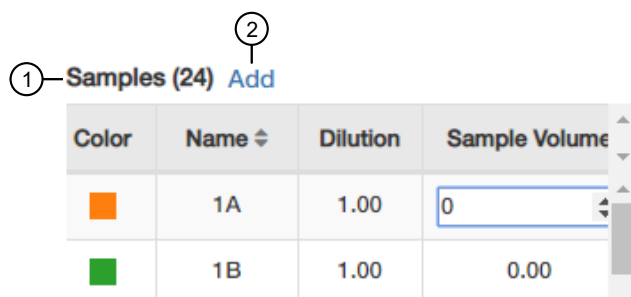


Figure 2 resDNASEQ™ template default cycling conditions

- Click **Next**.
- In the **Samples** pane of the **Setup** tab, enter the sample **Name** and **Dilution**. **Sample Volume** is not applicable. Add additional **Samples** if needed.

IMPORTANT! Do not change the **Targets**.



- Samples** pane
- Add**—adds additional samples

6. In the **Samples** pane of the **Setup** tab, scroll to the right, then enter the spike information.

For more information on plate setup, see the *AccuSEQ™ Real-Time PCR Software v3.1 User Guide* (Pub. No. 100094287).

- **Sample Volume**—not applicable; leave as default (0).
- **Spike Volume**—volume of DNA added to the PCR (set to 10).
- **Spike Standard Concentration**—expected spike amount per reaction (for example, 10 pg or 300 copies).
- **Reference**—the non-spiked sample; the mean quantity of reference is subtracted during % recovery calculation.
- **Spike Input**—automatically calculated (double check if correct).

Note: If incorrect, be sure **Spike Volume** is set to 10 and **Spike Standard Concentration** is the expected pg spike per PCR reaction.

- (Optional) **Comments**
- **Protein Concentration**—Drug substance protein concentration (if Total DNA in pg DNA/mg Protein is required).

Samples (24) Add


| | Spike Volume | Spike Standard Concentration |
|---|--------------|------------------------------|
| ① | 0 | 0.00 |
| | 0.00 | 0.00 |
| | 0.00 | 0.00 |
| | 0.00 | 0.00 |
| ② | | |

- ① Textbox—type in the value, or use the up and down arrows
 ② Scroll bar—scroll to find the spike parameter

7. Click **Next**.

The **Run** tab is displayed.

8. Experiments are auto-saved in the software. To save, exit the experiment. The software prompts you to save changes. Click **Yes**.

Note: Clicking  **Save As** will create a copy of the experiment.

Start the run

Start the run in the AccuSEQ™ Software.

| Option | Description |
|-----------------------------|---|
| If the experiment is open | Click Start Run . |
| If the experiment is closed | <ol style="list-style-type: none"> 1. Open the experiment. 2. Click the Run tab. 3. Click Start Run. |

The screenshot shows the AccuSEQ software interface. At the top, there is a navigation bar with tabs: Setup, Run, Result, Report, and Audit & E-Sign. The 'Run' tab is selected and highlighted with a circled '1'. Below the navigation bar, there are several icons for analysis settings, save, save as template, and export. The main area displays experiment details: Experiment Name: 2019-10-11_9189, E-Sign Status: Unsigned, and Assay Name: NewDNA_5Std. Below this, there are two sub-tabs: 'Plots' and 'Pool Run Summary'. The 'Plots' sub-tab is active, showing an 'Amplification Plot' on the left and a 'Well Plate' view on the right. The 'Amplification Plot' shows a y-axis labeled 'ΔF' ranging from 1e-1 to 1e+2 and an x-axis labeled 'Cycle' ranging from 5 to 60. The 'Well Plate' view shows a grid of wells (A-G, 1-11) with various sample types and cycle numbers. At the bottom right of the interface, there is a blue 'Start Run' button, which is circled with a '2'.

- ① Run tab
- ② Start Run button

A message stating **Run has been started successfully** is displayed when the run has started.

Analyze the results

After the qPCR run is finished, use the following general procedure to analyze the results. For more detailed instructions see the *AccuSEQ™ Real-Time PCR Software v3.1 User Guide* (Pub. No. 100094287).

1. In the AccuSEQ™ Real-Time PCR Software, open your experiment, then navigate to the **Result** tab.

- ① **Result** tab
- ② **Analysis Settings**
- ③ Plot horizontal scrollbar
- ④ **Analyze** button

2. In the **Result Analysis** tab, review the Amplification Curve plots for amplification profiles in the controls, samples, and the standard curve.
3. In the **Result Analysis** tab, review the **QC Summary** for any flags in wells.

4. In the **Result Analysis** tab, review the **Standard Curve** plot. Verify the values for the Slope, Y-intercept, R^2 , and Efficiency.

Note: The **Standard Curve** efficiency should be between 90-110% and the $R^2 > 0.99$. If these criteria are not met, up to two points, not in the same triplicate, can be removed from the standard curve data, and the analysis repeated.

5. (Optional) Navigate to the **Report** tab to generate a report of the experiment, or to export results.

3

Plasmid DNA quantitation

Sample residual plasmid DNA is measured in copy number, not concentration. To manually convert the copy number output to a mass measurement, multiply the copy number given by the AccuSEQ™ Real-Time PCR Software to the average molecular weight of your plasmid.

Prepare the control DNA serial dilutions for the standard curve

Guidelines for standard dilutions

- Prepare the standard curve and the test samples in different areas of the lab.
- Use different sets of pipettors for test sample preparation and for standard curve preparation and aliquoting to avoid cross-contamination of test samples.
- Vortex each tube for 15-30 seconds to mix the contents thoroughly before each dilution step.
- Vortex for 15-30 seconds, then tap down standards, before adding the standards to the PCR plate (plasmid standards are more fragile than genomic DNA standards).

Prepare the control serial dilutions (Plasmid DNA - Kanamycin Resistance)

1. Label nonstick 1.5-mL microfuge tubes: **NTC**, **SD1**, **SD2**, **SD3**, **SD4**, **SD5**, and **SD6**.
The dilution **SD6** will not be used for the standard curve. It will be used to confirm the limit of detection (LOD).
2. Add 70 µL of DNA Dilution Buffer (DDB) to tube NTC. Put aside.
3. Add 990 µL of DNA Dilution Buffer (DDB) to tube Dilution 1.
4. Add 180 µL of DNA Dilution Buffer (DDB) to tubes SD1, SD2, SD3, SD4, and SD5.
5. Add 100 µL of DNA Dilution Buffer (DDB) to tube SD6.
6. Remove the tube of KanR DNA control (3.0×10^7 copies/µL) from the freezer.
7. After the DNA thaws, vortex it thoroughly for 15-30 seconds, then tap gently to bring contents to the bottom of the tube. Do not centrifuge.
8. Perform the serial dilutions:
 - a. Add 10 µL of the KanR DNA control (3.0×10^7 copies/µL) to the tube that is labeled Dilution 1, vortex thoroughly for 15-30 seconds, then tap gently to bring contents to the bottom of the tube.

- b. Transfer 20 μL of the DNA from tube Dilution 1 to tube SD1, vortex thoroughly for 15-30 seconds, then tap gently to bring contents to the bottom of the tube..
- c. Continue to transfer 20 μL of DNA from the previous dilution tube to the next dilution tube until you add DNA to tube SD5.
- d. Transfer 100 μL of DNA from SD5 to SD6, then vortex thoroughly. Final dilutions are shown in the table. After each transfer, vortex thoroughly for 15-30 seconds, then tap gently to bring contents to the bottom of the tube. Do not centrifuge.

| Serial dilution (SD) tube | Dilution | Concentration (copy number/ μL) | Copy number/ PCR reaction |
|---------------------------|--|---|---------------------------|
| Control | DNA control tube | 3.0×10^7 | N/A |
| Dilution 1 | 10 μL DNA control + 990 μL DDB | 300,000 | N/A |
| SD 1 | 20 μL Dilution 1 + 180 μL DDB | 30,000 | 300,000 |
| SD 2 | 20 μL SD1 + 180 μL DDB | 3,000 | 30,000 |
| SD 3 | 20 μL SD2 + 180 μL DDB | 300 | 3,000 |
| SD 4 | 20 μL SD3 + 180 μL DDB | 30 | 300 |
| SD 5 | 20 μL SD4 + 180 μL DDB | 3 | 30 |
| SD 6 (LOD) | 100 μL SD5 + 100 μL DDB | 1.5 | 15 |

9. Store the DNA dilution tubes:

| Temperature | For use |
|-------------|---|
| 4°C | ≤ 2 days |
| -20°C | Dilution 1 ≤ 1 week |
| -20°C | Dilution 1 in single-use aliquots ≤ 6 months |

Prepare the samples

Extract samples with the PrepSEQ™ Residual DNA Sample Preparation Kit (Cat. No. 4413686).

1. Dilute the **Yeast tRNA**.

Table 11 Diluted Yeast tRNA

| Component | Volume |
|----------------------|-------------------------------------|
| Yeast tRNA (10mg/mL) | 5 μL |
| PBS (1X), pH 7.2 | 245 μL |
| Total | 250 μL |

2. Add 5 μL **Diluted Yeast tRNA** to 370 μL of each test sample. This is sufficient for triplicate 100 μL extractions.

Prepare the PCR reaction mix

1. Determine the number of reactions needed for the controls and test samples that you will quantify.
2. Thaw all kit reagents completely at room temperature, thoroughly mix reagents, and briefly centrifuge.
3. Prepare a PCR reaction mix using the reagents and volumes shown in the table below.
 - Multiply the PCR reaction volume for one reaction (30 μ L) by the number of reactions that you need to run.
 - Use 10% excess volume to compensate for pipetting losses.

Note: Use reagents from the same lot for all reactions.

| Kit reagents | Volume for 1 30- μ L reaction | Volume for 48 30- μ L reactions (includes 10% overage) |
|--|-----------------------------------|---|
| Negative Control (water) | 2 μ L | 106 μ L |
| 10X DNA assay mix appropriate for the cell line being tested | 3 μ L | 159 μ L |
| 2X Environmental Master Mix | 15 μ L | 795 μ L |
| DNA template | 10 μ L | Add DNA template to each well separately, not as part of Master Mix |
| Total | 30 μL | 1060 μL |

Prepare the PCR plate

Plate setup differs slightly for each AccuSEQ™ System. See your software user guide for specific instructions. Place samples, NTCs, and standards in different quadrants of the plate.

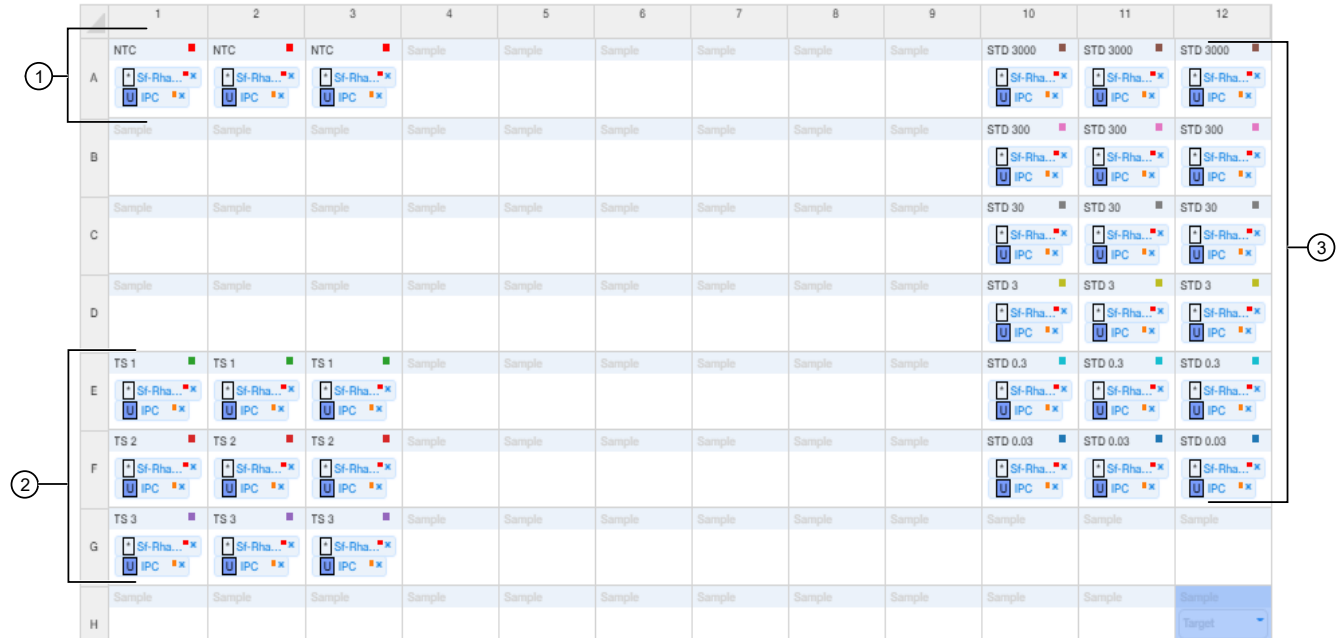


Figure 3 Default plate setup in the AccuSEQ™ Real-Time PCR Software v3.1

- ① No template controls
- ② Samples
- ③ Standard curve

1. Add 20 µL PCR reaction mix to each well.
2. Add 10 µL of PCR NTC to the appropriate wells.
3. Add 10 µL each of sample DNA (with tRNA added) to the appropriate wells.

IMPORTANT! Vortex samples for 15-30 seconds before adding the samples to the **PCR Plate**, then tap to bring contents to the bottom of the wells.

If you prepared samples with the automated protocol, use a multichannel pipette to transfer the extracted sample.

4. Add 10 µL of standard dilutions to the appropriate wells.

IMPORTANT! Vortex samples for 15-30 seconds before adding the samples to the **PCR Plate**, then tap to bring contents to the bottom of the wells.

If you prepared samples with the automated protocol, use a multichannel pipette to transfer the extracted sample.

5. Use a film applicator to seal the plate with optical film, then briefly centrifuge with a miniplate centrifuge that is compatible with 96-well plates.

Setup, run, and analyze samples with AccuSEQ™ Software on the QuantStudio™ 5 Real-Time PCR Instrument

Create a resDNASEQ™ template (Plasmid DNA)

Plasmid DNA resDNASEQ™ assays do not use a **Factory default/Admin Defined Template**. Create a new template before the first use of these assays.

1. In the **Home** screen, click **Templates** in the left navigation pane.

| Template Name | Template Location | Publish | Created Date | Created By User | Modified Date | Modified By User | Actions |
|--------------------------------|-------------------|---------|-------------------------------|-----------------|-------------------------------|------------------|---------|
| 2018-11-14_ProA 2020-06-03-... | My Template | | 03 Jun 2020 20:01:23 GMT-0500 | Administrator | 03 Jun 2020 20:01:46 GMT-0500 | Administrator | + ✎ 📄 🔒 |
| Mycoseq | Factory Default | ☑ | 01 Jun 2020 06:44:57 GMT-0500 | system | 01 Jun 2020 06:44:57 GMT-0500 | system | + ✎ 📄 🔒 |
| ResDNA_5Std | Factory Default | ☑ | 01 Jun 2020 06:44:57 GMT-0500 | system | 01 Jun 2020 06:44:57 GMT-0500 | system | + ✎ 📄 🔒 |
| ProteinSeq HCP | Factory Default | ☑ | 01 Jun 2020 06:44:57 GMT-0500 | system | 01 Jun 2020 06:44:57 GMT-0500 | system | + ✎ 📄 🔒 |
| ProteinSeq Pro A | Factory Default | ☑ | 01 Jun 2020 06:44:57 GMT-0500 | system | 01 Jun 2020 06:44:57 GMT-0500 | system | + ✎ 📄 🔒 |
| ResDNA_6Std | Factory Default | ☑ | 01 Jun 2020 06:44:57 GMT-0500 | system | 01 Jun 2020 06:44:57 GMT-0500 | system | + ✎ 📄 🔒 |

1. **Templates** icon

2. Click **+ Create New** next to the **ResDNA_5Std** factory default template.
3. Click **Next** to move to the **qPCR Method** screen.
4. Click **Next** to move to the **Plate Setup** screen.
5. In the **Plate Setup** screen, add the **Targets** and **Reporters**.

| <input type="checkbox"/> | Color | Name | Reporter | Quencher | Task | Qua |
|--------------------------|-------|------|----------|----------|------|-----|
| <input type="checkbox"/> | ■ | Kan | FAM | NFQ-MGB | | |
| <input type="checkbox"/> | ■ | IPC | NED | NFQ-MGB | | |

For the resDNASEQ™ Quantitative Plasmid DNA - Kanamycin Resistance Gene Kit, this is the FAM™ dye for the kanamycin (Kan) target and the NED™ dye for the IPC.

6. Click **Save as Template**.
7. Enter a **Template Name** and description, then select **Admin Defined** and **Locked**. Click **Save**. The template is saved, and can be accessed from **Templates** in the **Home** screen.
8. Click **Templates** in the **Home** screen, then open the new template.

Note: The template must be saved prior to editing the **Analysis Settings**




9. Click  **Analysis Settings**, then deselect **Default Settings**.
10. Enter new thresholds for the targets.


Table 12 resDNASEQ™ Quantitative Plasmid DNA - Kanamycin Resistance Gene Kit

| Target | Threshold |
|-----------|-----------|
| Kanamycin | 0.04 |
| IPC | 0.02 |

11. Click **Apply**, then close the template.
12. In the  **Home** screen, click  **Templates** in the left navigation pane, then **Publish** the template. See *AccuSEQ™ Real-Time PCR Software v3.1 User Guide* (Pub. No. 100094287). The template is listed in the **Factory default/Admin Defined Templates**.

Create a resDNASEQ™ experiment (Plasmid DNA)

Plasmid DNA resDNASEQ™ assays do not use a **Factory default/Admin Defined Template**. Create a new template before the first use of these assays.

1. In the  **Home** screen, click **Factory default/Admin Defined Templates**, then select the custom plasmid DNA **resDNASEQ** template created in “Create a resDNASEQ™ template (Plasmid DNA)” on page 28.

Note: To create an experiment from an existing resDNASEQ™ experiment, see *AccuSEQ™ Real-Time PCR Software v3.1 User Guide* (Pub. No. 100094287).

2. In the **Experiment Properties** pane of the **Setup** tab:
 - a. (Optional) Change the system-generated name of the experiment.

Note: Names must be unique. Deleted experiment names can not be reused.

- b. (Optional) Enter the plate **Barcode**, then add **Comments**.

Note: Comments are not editable post analysis.

Default resDNASEQ™ settings (cannot be changed)

- **Experiment Type** is **Quantitation-Standard Curve**
- **Chemistry** is **TaqMan™ Reagents**
- **Ramp Speed** is **Standard - 2hrs**

- c. Click **Next**.

Note: Experiment names cannot be changed after this step.

- In the **qPCR Method** pane of the **Setup** tab, view the default volume and cycling conditions (cannot be changed).

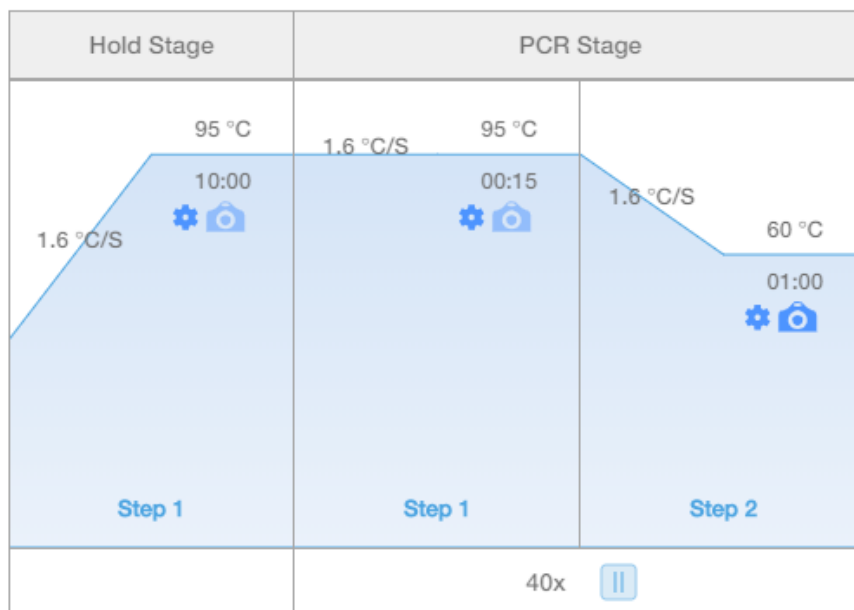


Figure 4 resDNASEQ™ template default cycling conditions

- Click **Next**.
- In the **Samples** pane of the **Setup** tab, enter the sample **Name** and **Dilution**. **Sample Volume** is not applicable. Add additional **Samples** if needed.

IMPORTANT! Do not change the **Targets**.

① — Samples (24) Add

| Color | Name ↕ | Dilution | Sample Volume |
|--------|--------|----------|---------------|
| Orange | 1A | 1.00 | 0 |
| Green | 1B | 1.00 | 0.00 |

- Samples** pane
- Add**—adds additional samples

6. In the **Samples** pane of the **Setup** tab, scroll to the right, then enter the spike information.

For more information on plate setup, see *AccuSEQ™ Real-Time PCR Software v3.1 User Guide* (Pub. No. 100094287).

- **Sample Volume**—not applicable; leave as default (0).
- **Spike Volume**—volume of DNA added to the PCR (set to 10).
- **Spike Standard Concentration**—expected spike amount per reaction.
- **Reference**—the non-spiked sample; the mean quantity of reference is subtracted during % recovery calculation.
- **Spike Input**—automatically calculated (double check if correct).

Note: If incorrect, be sure **Spike Volume** is set to 10 and **Spike Standard Concentration** is the expected pg spike per PCR reaction.

- (Optional) **Comments**
- **Protein Concentration**—Drug substance protein concentration (if Total DNA in pg DNA/mg Protein is required).

Samples (24) Add

| | Spike Volume | Spike Standard Concentration |
|---|--------------|------------------------------|
| ① | 0 | 0.00 |
| | 0.00 | 0.00 |
| | 0.00 | 0.00 |
| | 0.00 | 0.00 |
| ② | | |

① Textbox—type in the value, or use the up and down arrows

② Scroll bar—scroll to find the spike parameter

For more information on plate setup, see *AccuSEQ™ Real-Time PCR Software v3.1 User Guide* (Pub. No. 100094287).

The screenshot displays the software interface for setting up a PCR plate. At the top, the 'E-Sign Status' is 'Unsigned' and the 'Assay Name' is 'ResDNA_5Std'. A toolbar (1) contains icons for Undo, Redo, Copy, Paste, Delete, and View. Below the toolbar are two dropdown menus for 'Select well:' and 'Select:' with a 'Define & setup Standard' button. The main area is a 96-well plate grid (rows A-H, columns 1-12). Row A contains 3 NTC wells (2) and 3 Sample wells. Row B contains 3 Sample wells. Row C contains 3 Sample wells. Row D contains 3 Standard wells (4) with concentrations of 3000.0000, 300.0000, and 30.0000. Row E contains 3 TS 1 wells (3), Row F contains 3 TS 2 wells (3), and Row G contains 3 TS 3 wells (3). The bottom status bar shows 'Wells: U 27 N 3 S 15 69 (Empty)' and a progress indicator at 0%.

Figure 5 resDNASEQ™_5Std template default sample plate layout

- ① Toolbar (in order: Undo, Redo, Copy, Paste, Delete, View)
- ② 3 No Template Control (NTC) samples
- ③ 3 default **Samples**
- ④ Standard curve dilutions (S) in triplicate

7. Click **Next**.

The **Run** tab is displayed.

8. Experiments are auto-saved in the software. To save, exit the experiment. The software prompts you to save changes. Click **Yes**.

Note: Clicking **Save As** will create a copy of the experiment.

9. Assemble the PCR reactions following the manufacturer's instructions for the reagents and following the plate layout set up in the template.

Start the run

Start the run in the AccuSEQ™ Software.

| Option | Description |
|-----------------------------|---|
| If the experiment is open | Click Start Run . |
| If the experiment is closed | <ol style="list-style-type: none"> 1. Open the experiment. 2. Click the Run tab. 3. Click Start Run. |

The screenshot shows the AccuSEQ software interface. At the top, there is a navigation bar with tabs: Setup, Run, Result, Report, and Audit & E-Sign. The 'Run' tab is selected and highlighted with a circled '1'. Below the navigation bar, there are buttons for Analysis Settings, Save As, Save As Template, and Export Experiment. The main area displays 'Run Control' with sub-tabs for Plots and Pool Run Summary. Under 'Plots', 'Amplification' is selected. An 'Amplification Plot' is shown on the left, and a 'Well Plate' view is on the right. At the bottom right, there is a 'Start Run' button highlighted with a circled '2'.

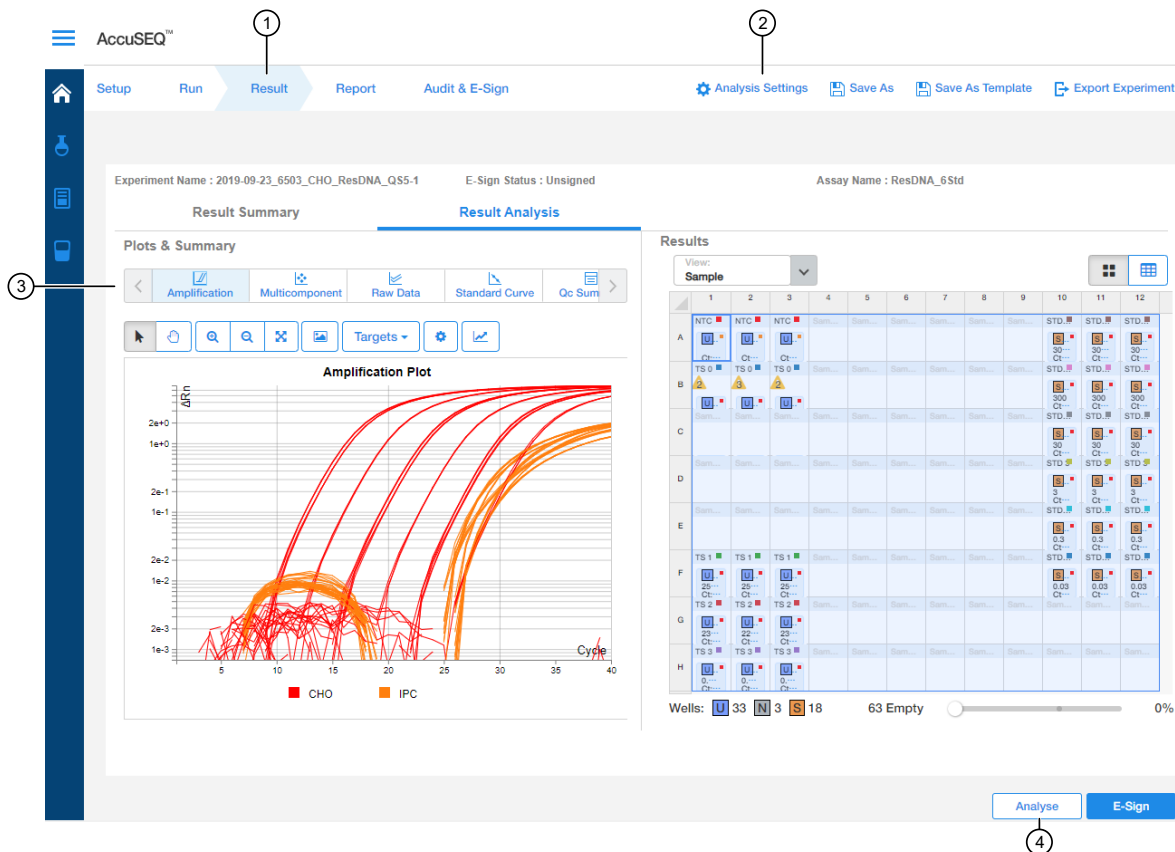
- ① Run tab
- ② Start Run button

A message stating **Run has been started successfully** is displayed when the run has started.

Analyze the results

After the qPCR run is finished, use the following general procedure to analyze the results. For more detailed instructions see the *AccuSEQ™ Real-Time PCR Software v3.1 User Guide* (Pub. No. 100094287).

1. In the AccuSEQ™ Real-Time PCR Software, open your experiment, then navigate to the **Result** tab.



- ① **Result** tab
- ② **Analysis Settings**
- ③ Plot horizontal scrollbar
- ④ **Analyse** button

2. In the **Result Analysis** tab, review the Amplification Curve plots for amplification profiles in the controls, samples, and the standard curve.

Note: Ensure that the threshold was set to 0.04 for Kanamycin target and 0.02 for IPC.

3. In the **Result Analysis** tab, review the **QC Summary** for any flags in wells.

4. In the **Result Analysis** tab, review the **Standard Curve** plot. Verify the values for the Slope, Y-intercept, R^2 , and Efficiency.

Note: The **Standard Curve** efficiency should be between 90-110% and the $R^2 > 0.99$. If these criteria are not met, up to two points, not in the same triplicate, can be removed from the standard curve data, and the analysis repeated.

5. (Optional) Navigate to the **Report** tab to generate a report of the experiment, or to export results.
6. (Optional) To manually convert the copy number output to a mass measurement, multiply the copy number given by the AccuSEQ™ Real-Time PCR Software to the average molecular weight of your plasmid.



Troubleshooting

| Observation | Possible cause | Action |
|--|--|---|
| <p>Slope for the standard curve is outside the typical range, or R_2 value is significantly less than 0.99.</p> | <p>When applying detectors for standards, the Task and Quantity were applied to the wrong detector.</p> <p><i>or</i></p> <p>The incorrect Quantity was entered.</p> <p><i>or</i></p> <p>Adjust baseline settings.</p> <p><i>or</i></p> <p>Poor standard curve preparation technique (forgot to mix, inaccurate pipetting).</p> | <ol style="list-style-type: none"> 1. In the SDS software, from the plate document, double-click a well containing a DNA standard to view the Well Inspector. 2. Ensure that the correct Task and Quantity are applied to the correct detector, then reanalyze. 3. Compare std curve statistics using autobaseline or manual baseline. The upper limit of the manual baseline setting must be 2 cycles before uptick in amplification. Verify in R_n vs C_t linear view. |
| <p>ΔR_n and C_t values are inconsistent with replicates</p> | <p>Evaporation of reaction mixture from some wells occurred because the optical adhesive cover was not correctly sealed to the reaction plate or due to over-drying the eluates in PrepSEQ™.</p> | <ol style="list-style-type: none"> 1. Select the Component tab. Confirm that affected wells generated significantly less fluorescence than unaffected replicates. 2. Check the amount of solution in each well of the reaction plate. Confirm that the wells affected by evaporation contained less solution than unaffected wells, and corresponded with the inconsistent results. 3. For subsequent runs, ensure that the optical adhesive cover is correctly sealed to the reaction plate. |

(continued)

| Observation | Possible cause | Action |
|---|--|--|
| ΔR_n and C_t values are inconsistent with replicates | Incorrect volume of PCR reaction mix was added to some reactions. | <ol style="list-style-type: none"> 1. Select the Component tab. Confirm that affected wells generated significantly less fluorescence than unaffected replicates. 2. Select the Spectra tab. Confirm that the wells with the incorrect volume of PCR reaction mix generated significantly different amounts of fluorescence than the unaffected wells. 3. For subsequent runs, ensure the correct volume of PCR reaction mix. |
| Jagged amplification plots | Weak lamp or incorrect replacement. | Replace the lamp or make sure that the existing replacement is correct. |
| No defined amplification plots | <p>An incorrect detector was selected on the amplification plot.</p> <p>or</p> <p>An incorrect detector was applied to the reactions when setting up the plate document.</p> | <ol style="list-style-type: none"> 1. Confirm that the correct detector was selected on the amplification plot. 2. If the correct detector was not selected, then in the plate document, double-click a well to view the Well Inspector, verify that the detector settings are correct, and reanalyze. |
| Abnormal ΔR_n values or negative ΔR_n values. | Incorrect passive reference was selected when setting up the plate document. | <ol style="list-style-type: none"> 1. From the plate document, double-click a well to view the Well Inspector. 2. Ensure that ROX™ was selected as the Passive Reference. |
| Standard curve for plasmid DNA assays is outside of the 90–110% efficiency range. | Incomplete vortexing of low level standards. | Repeat reactions, ensuring that samples and standards are vortexed for 15-30 seconds. |
| Wide variance of C_t values of plasmid DNA samples. | Incomplete vortexing of samples. | |



Use the kit with the 7500 Fast Real-Time PCR Instrument and AccuSEQ™ software v2.x

Required materials not supplied

Unless otherwise indicated, all materials are available through thermofisher.com. "MLS" indicates that the material is available from fisherscientific.com or another major laboratory supplier.

| Item | Source |
|---|---|
| Instrument | |
| 7500 Fast Real-Time PCR System with AccuSEQ™ software v2.x | Contact your local sales representative |
| Generic consumables | |
| Disposable gloves | MLS |
| Aerosol-resistant pipette tips | MLS |
| Pipettors: <ul style="list-style-type: none">• Positive-displacement• Air-displacement• Multichannel | MLS |
| Consumables for the 7500 Fast Real-Time PCR System | |
| MicroAmp™ Optical 96-Well Reaction Plate with Barcode, 20 plates, 0.2-mL well; for use with 7300, 7500, and 7900HT Fast Real-Time PCR Systems | 4306737^[1] |
| MicroAmp™ Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL, 20 plates; for use with 7500 Fast Real-Time PCR System | 4346906 |
| MicroAmp™ Optical 96-Well Reaction Plate with Barcode & Optical Adhesive Films, 100 plates with covers; for use with 7300 and 7500 Fast Real-Time PCR Systems | 4314320 |
| MicroAmp™ Optical 8-Cap Strips, 300 strips | 4323032 |

(continued)

| Item | Source |
|---|-------------------------|
| MicroAmp™ Optical Adhesive Film Kit, 20 covers, 1 applicator, 1 optical cover compression pad | 4313663 |
| MicroAmp™ Optical Adhesive Film | 4360954 |

^[1] Not recommended for use with the 7500 Fast system. For 7500 Fast system reactions, use Cat. No. [4346906](#).

Create a plate document in the AccuSEQ™ software

1. In the home screen, select **Create Custom Experiment**.
2. In the **Experiment name** field, enter a unique name for the experiment.
3. Specify experiment information.
 - a. Select experiment type **Quantitation – Standard Curve**.
 - b. Select reagents **TaqMan™ Reagents**.
 - c. Select ramp speed **Standard**.
4. In the **Plate Setup** screen, select the **Define Targets and Samples** tab.
5. Specify target information.
 - a. Click **Add New Target**.
 - b. Enter a host cell name in the target name field.
 - c. Select reporter **FAM™** and quencher **NFQ_MGB**.

Note: For the Sf9 and Baculovirus assay, select reporter VIC™ for the Sf9 target and quencher **NFQ_MGB**. Add an additional target, and select reporter FAM™ for the baculovirus target and quencher **NFQ_MGB**.

- d. Select a color for this target.
6. Specify IPC target information.
 - a. Click **Add New Target**.
 - b. Enter **IPC** in the target name field.
 - c. Select reporter **VIC™** and quencher **NFQ_MGB**.

Note: For the Sf9 and Baculovirus assay and the Plasmid DNA - Kanamycin Resistance assay, select the reporter NED™ and quencher NFQ_MGB.

- d. Select a color for this target.

7. Define new samples.
 - a. Click **Add New Sample**.
 - b. In **Sample Name**, add the names of the samples you want to define.
 - c. Click **Next**, or select the **Assign Targets and Samples** tab.
8. In the **Assign Targets and Samples** tab, define new targets.
 - a. Follow the instructions in the top of the tab to set up the standards, unknowns, and negative controls.
 - b. Click **Define and Set Up Standards** to open the **Define and Set Up Standards** dialog box to enter the appropriate settings and define the standard curve. When defined, click **Apply** and **Close**. The new standard curve is applied to the plate layout screen.
9. Assign the IPC to the standard curve wells.
10. In the **Run Method** screen, in the **Graphical View** tab.
 - a. In **Reaction Volume Per Well**, enter **30 µL**
 - b. Right-click the left column named **Holding Stage 1** and select **Delete Selected**. This 50°C hold stage is not needed.
11. Click the **Analysis** button in the left panel. In the **Analysis Settings** window on the right, set the default settings.
 - a. On the **Ct Settings** tab, click **Edit Default Settings**. Then set **Threshold** to 0.2, set to **Automatic Baseline**, and then click **Save Changes**.

Note: For CHO samples, a manual baseline of 3–12 is more appropriate.
For the Plasmid DNA - Kanamycin Resistance assay, do not use the default settings. Set the threshold to 0.04 for KanR and 0.02 for IPC.

 - b. Select (highlight) both targets.
 - c. In the right-hand window, select **Use Default Settings**.
 - d. Click **Apply Analysis Settings**.

12. Select **File ▶ Save as**, confirm that the file is named “resDNA_Template”, then select **Save as a template file** in the drop-down list and close the template plate document.

Note: You can reuse the plate template document to run an assay by opening the template file and choosing **Save As** to save the file with the experiment name.

Run the plate


1. In the toolbar, select **File** ▶ **Open**, navigate to the **resDNA_Template** file, then click **Open**.
2. In the **Experiment Name** field, enter the appropriate experiment name, then click **Finish**.
3. Make any necessary changes to the test sample labels.
 - **Sample Volume**—not applicable; leave as default (0).
 - **Spike Volume**—volume of DNA added to the PCR (set to 10).
 - **Spike Standard Concentration**—expected spike amount per reaction (for example, 10pg).
 - **Reference**—the non-spiked sample; the mean quantity of reference is subtracted during % recovery calculation.
 - **Spike Input**—automatically calculated (double check if correct).

Note: If incorrect, be sure **Spike Volume** is set to 10 and **Spike Standard Concentration** is the expected pg spike per PCR reaction.

4. Select **Save As** to save the new experiment as an EDS experiment file with the same name as entered in the **Experiment Name** field.
5. Load the plate into the instrument.
6. Click **Start Run**.
7. Select a run screen (**Amplification** plot, **Temperature** plot, or **Run method**) to monitor the progress of the run.

Analyze the results

After the qPCR run is finished, use the following general procedure to analyze the results:

1. In the toolbar, select **Analysis** ▶ **Analysis Settings**.
2. Click  (**Analyze**).
3. Select **Analysis** ▶ **QC Summary** in the left panel of the screen. Review the flag summary.
4. In the left panel, select **Analysis** ▶ **Standard Curve**. Verify the values for the Slope, Y-Intercept, R2, and Efficiency.
5. Select **File** ▶ **Print Report** to generate a hardcopy of the experiment, or click **Print Preview** to view and save the report as a *.pdf or *.html file.
6. Optional: Select **File** ▶ **Export**. In the **Export Data** menu, select file type *.xls. Click **Start Export**.



Use the kit with 7500 System SDS Software v1.5.1

Create the plate document, run the plate, and analyze the results with 7500 Fast SDS software

The following instructions apply only to the Applied Biosystems™ 7500 Fast instrument with SDS v1.x software. If you use a different instrument or software, refer to the applicable instrument or software documentation.

Create a plate document

Residual DNA assays are duplex assays, containing sample DNA and Internal Positive Control (IPC).

Plate document: settings

If you have run the assay frequently, you can use the table below to enter settings in Plate Document fields. If you are a new user, follow the detailed instructions in the following sections.

| Summary of settings for the Plate Document | | |
|--|----------------------------------|--|
| In this field... | | Use these settings |
| Detector | resDNASEQ™ kit target cell lines | <ul style="list-style-type: none">• Single target assays: FAM™• Sf9 and Baculovirus assay: VIC™ and FAM™ (Select NFQ_MGB for quencher) |
| | IPC | <ul style="list-style-type: none">• Single target assays: VIC™• Sf9 and Baculovirus assay and Plasmid DNA - Kanamycin Resistance assay: NED™ (Select NFQ_MGB for quencher) |
| PCR | Hold | Temp: 95°C Time: 10:00 |



(continued)

| Summary of settings for the Plate Document | | |
|--|---|---|
| In this field... | | Use these settings |
| PCR | Cycling (Standard Mode) | Cycles: 40 Temp: 95°C Time: 0:15 Temp: 60°C Time: 1:00 |
| Analysis | CHO, <i>E. coli</i> , HEK293, Human, MDCK, NS0, <i>Pichia</i> , Plasmid DNA - Kanamycin resistance, Sf9 and Baculovirus, and Vero | Automatic Baseline or Manual Baseline ^[1] Threshold: 0.2 Note: For CHO, the upper limit for manual baseline analysis is 12. For the Plasmid DNA - Kanamycin Resistance assay, change the threshold to 0.04 for KanR and 0.02 for IPC. |

^[1] You can analyze the assay using Automatic or Manual Baseline, use the setting that yields the best standard curve.

Plate document: procedure

In the SDS software:

1. In the template **Assay** drop-down list, select **Absolute Quantification**.
2. In the **Run Mode** drop-down list, select **Standard 7500**.
3. Enter **resDNA_Template** in the **Plate** name field, then click **Next**.
4. Click **New Detector**:
 - a. Enter the name of the target cell line in the **Name** field.
 - b. Select reporter **FAM™** and quencher **NFQ_MGB**.

Note: For the Sf9 and Baculovirus assay, select reporter **VIC™** for the Sf9 target and quencher **NFQ_MGB**. Add an additional target, and select reporter **FAM™** for the baculovirus target and quencher **NFQ_MGB**.

- c. Select a color for the detector, then click **Create Another**.
5. Click **New Detector**:
 - a. Enter **IPC** in the **Name** field.
 - b. Select reporter **VIC™** and quencher **NFQ_MGB**.

Note: For the Sf9 and Baculovirus assay and the Plasmid DNA - Kanamycin Resistance assay, select the reporter **NED™** and the quencher **NFQ_MGB**.

 - c. Select a color for the detector, then click **OK**.
 - d. Select the detectors, then click **Add>>** to add the detectors to the document (plate).



6. Select **ROX™** as the passive reference dye, then click **Next**.
7. Select the applicable set of wells for the samples, then select the target cell line and **IPC** detectors for each well. The following figure shows an example plate layout:

| | | | | | | | | | | Standard Curve (pg) | | |
|---|-----|-----|-----|---|------------|------------|------------|---|---|---------------------|----------|----------|
| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
| A | TS1 | TS1 | TS1 | | TS1 ERC | TS1 ERC | TS1 ERC | | | NTC | NTC | NTC |
| B | TS2 | TS2 | TS2 | | TS2 ERC | TS2 ERC | TS2 ERC | | | | | |
| C | TS3 | TS3 | TS3 | | TS3 ERC | TS3 ERC | TS3 ERC | | | 0.03 pg | 0.03 pg | 0.03 pg |
| D | | | | | | | | | | 0.3 pg | 0.3 pg | 0.3 pg |
| E | | | | | | | | | | 3 pg | 3 pg | 3 pg |
| F | NEG | NEG | NEG | | | | | | | 30 pg | 30 pg | 30 pg |
| G | | | | | | | | | | 300 pg | 300 pg | 300 pg |
| H | | | | | | | | | | 3,000 pg | 3,000 pg | 3,000 pg |

8. Set tasks for each sample type by clicking on the **Task Column** drop-down list:
 - a. NTC: target cell line detector task = **NTC**
 - b. NEG, test samples, and ERC wells: target DNA detector task = **Unknown**
 - c. IPC = **Unknown** for all wells
9. Set up the standard curve:
 - a. Select the wells.
 - b. Assign the tasks (target DNA = **Standard**) and enter the appropriate Quantity for each set of triplicates.

| Tube label | Row-wells | Task | Quantity | Label (pg) |
|------------|--------------|----------|----------|------------|
| SD 1 | H-10, 11, 12 | Standard | 3,000 | 3,000 pg |
| SD 2 | G-10, 11, 12 | Standard | 300 | 300 pg |
| SD 3 | F-10, 11, 12 | Standard | 30 | 30 pg |
| SD 4 | E-10, 11, 12 | Standard | 3 | 3 pg |



(continued)

| Tube label | Row-wells | Task | Quantity | Label (pg) |
|--|--------------|----------|----------|------------|
| SD 5 | D-10, 11, 12 | Standard | 0.3 | 0.3 pg |
| SD 6 (for CHO, Vero, MDCK, and NS0 only) | C-10, 11, 12 | Standard | 0.03 | 0.03 pg |

Note: The Plasmid DNA - Kanamycin Resistance assay creates a standard curve based on copy number (300,000 to 30 copies per reaction).

10. Select the **Instrument** tab, then set thermal-cycling conditions:

- Set the **thermal cycling reaction volume** to **30 µL**.
- Set the reaction to **Standard Mode**.
- Set the temperature and the time as shown in the following table:

| Step | AmpliAq Gold™ enzyme activation | PCR | |
|---------------|---------------------------------|-------------------|---------------|
| | | Denature | Anneal/extend |
| | Hold | Cycle (40 Cycles) | |
| Temp (°C) | 95 | 95 | 60 |
| Time (mm:sec) | 10:00 | 0:15 | 1:00 |

Refer to the applicable 7500 Fast Real-Time PCR Systems instrument manual for additional information.

11. In the **Analysis Settings** window, enter the following settings, then click **OK**:

- a. Select **Manual Ct**.
- b. In **Threshold**, enter **0.2**.

Note: For the Plasmid DNA - Kanamycin Resistance assay, change the threshold to 0.04 for KanR and 0.02 for IPC.

- c. Select **Automatic Baseline** or **Manual Baseline**.

Note: You can analyze the assays using Automatic or Manual Baseline, use the setting that yields the best standard curve. For CHO, the upper limit threshold for manual baseline analysis is 12.

12. Select **File ▶ Save as**, confirm that the file is named “resDNA_Template”, then select **SDS Templates (*.sdt)** in the **Save as type** drop-down list and close the template plate document.

Note: You can reuse the plate template document whenever you run the assay.

13. Close the saved template file.



Run the plate

1. In the SDS software, select **File ▶ New**, navigate to the **resDNA_Template** file (created in “Plate document: procedure” on page 43), then click **Open**.
2. In **Plate Name**, enter an appropriate experiment name, then click **Finish**.
3. Make any necessary changes to the test sample labels.
4. Select **Save As** to save the new experiment as an SDS experiment file.
5. Load the plate on the instrument.
6. Select the **Instrument** tab, save the document, then click **Start** to start the real-time qPCR run.

Analyze the results

After the qPCR run is finished, use the following general procedure to analyze the results.

1. Select the **Results** tab.
2. Select the **Amplification Plot** tab.
3. Verify the analysis settings, change as appropriate, then click **Analyze**.
4. Select the **Results tab ▶ Standard Curve** tab, then verify the Slope, Intercept, and R2 values.
5. Right-click the **Standard Curve**, select **Export as JPEG**, then click **OK**. Alternatively, press **PrintScreen**, then paste the image in a WordPad file.
6. Select the **Report tab ▶ Report**, then review the mean quantity and standard deviation for each sample.
7. Optional: Select **File ▶ Export ▶ Results**. In the **Save as type** drop-down list, select **Results Export Files (*.csv)**, then click **Save**.



Good laboratory practices

Work area setup and lab design

The sensitivity of this kit (and other PCR-based tests) enables amplification of minute quantities of DNA, necessitating precautions to avoid contamination of samples yet to be amplified (Kwok and Higuchi, 1989).

Process samples carefully to prevent contamination by human DNA. Wear gloves at all times and change them frequently. Close sample tubes when not in use. Limit aerosol dispersal by handling sample tubes and reagents carefully.

Good laboratory practices for PCR and RT-PCR

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

Avoiding false positives due to cross-contamination

To avoid false positives due to cross-contamination:

- Do not open tubes after amplification.
- Use different sets of pipettors when pipetting negative control, unknown, and positive control samples.

Note: Refer to “Prepare the PCR plate” on page 16 for best practice.



Safety



WARNING! GENERAL SAFETY. Using this product in a manner not specified in the user documentation may result in personal injury or damage to the instrument or device. Ensure that anyone using this product has received instructions in general safety practices for laboratories and the safety information provided in this document.

- Before using an instrument or device, read and understand the safety information provided in the user documentation provided by the manufacturer of the instrument or device.
- Before handling chemicals, read and understand all applicable Safety Data Sheets (SDSs) and use appropriate personal protective equipment (gloves, gowns, eye protection, and so on). To obtain SDSs, see the “Documentation and Support” section in this document.

Chemical safety



WARNING! GENERAL CHEMICAL HANDLING. To minimize hazards, ensure laboratory personnel read and practice the general safety guidelines for chemical usage, storage, and waste provided below. Consult the relevant SDS for specific precautions and instructions:

- Read and understand the Safety Data Sheets (SDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials. To obtain SDSs, see the "Documentation and Support" section in this document.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing).
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with sufficient ventilation (for example, fume hood).
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer cleanup procedures as recommended in the SDS.
- Handle chemical wastes in a fume hood.
- Ensure use of primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container. Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- After emptying a waste container, seal it with the cap provided.
- Characterize (by analysis if needed) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure that the waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.
- **IMPORTANT!** Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.



WARNING! HAZARDOUS WASTE (from instruments). Waste produced by the instrument is potentially hazardous. Follow the guidelines noted in the preceding General Chemical Handling warning.



WARNING! 4L Reagent and Waste Bottle Safety. Four-liter reagent and waste bottles can crack and leak. Each 4-liter bottle should be secured in a low-density polyethylene safety container with the cover fastened and the handles locked in the upright position.

Biological hazard safety



WARNING! Potential Biohazard. Depending on the samples used on this instrument, the surface may be considered a biohazard. Use appropriate decontamination methods when working with biohazards.



WARNING! BIOHAZARD. Biological samples such as tissues, body fluids, infectious agents, and blood of humans and other animals have the potential to transmit infectious diseases. Conduct all work in properly equipped facilities with the appropriate safety equipment (for example, physical containment devices). Safety equipment can also include items for personal protection, such as gloves, coats, gowns, shoe covers, boots, respirators, face shields, safety glasses, or goggles. Individuals should be trained according to applicable regulatory and company/ institution requirements before working with potentially biohazardous materials. Follow all applicable local, state/provincial, and/or national regulations. The following references provide general guidelines when handling biological samples in laboratory environment.

- U.S. Department of Health and Human Services, *Biosafety in Microbiological and Biomedical Laboratories (BMBL)*, 5th Edition, HHS Publication No. (CDC) 21-1112, Revised December 2009; found at:
<https://www.cdc.gov/labs/pdf/CDC-BiosafetymicrobiologicalBiomedicalLaboratories-2009-P.pdf>
- World Health Organization, *Laboratory Biosafety Manual*, 3rd Edition, WHO/CDS/CSR/LYO/2004.11; found at:
www.who.int/csr/resources/publications/biosafety/Biosafety7.pdf

Documentation and support

Related documentation

| Document | Publication number | Description |
|---|--------------------|--|
| <i>resDNASEQ™ Quantitative DNA Kits Quick Reference</i> | 4469837 | For brief instructions on using the resDNASEQ™ Quantitative DNA Kits. |
| <i>PrepSEQ™ Sample Preparation Kits User Guide</i> | 4469838 | For information on preparing samples for extraction. |
| <i>PrepSEQ™ Residual DNA Sample Preparation Kit Quick Reference</i> | 4469839 | For brief instructions on preparing samples for extraction. |
| <i>AccuSEQ™ Real-Time PCR Software v3.1 User Guide</i> | 100094287 | For information on AccuSEQ™ Real-Time PCR Software v3.1 with the QuantStudio™ 5 Real-Time PCR System |
| <i>AccuSEQ™ Real-Time PCR Software v3.1 Quick Reference</i> | 100094288 | For basic information on AccuSEQ™ Real-Time PCR Software v3.1 with the QuantStudio™ 5 Real-Time PCR System |
| <i>Applied Biosystems™ 7300/7500/7500 Fast Real-Time PCR System Getting Started Guide: Absolute Quantitation using Standard Curve</i> | 4347825 | For information on the 7500 Fast instrument. |
| <i>AccuSEQ™ software: Custom Quick Reference Card</i> | 4425585 | For information on AccuSEQ™ software with the 7500 Fast instrument. |

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- Product documentation
 - User guides, manuals, and protocols
 - Certificates of Analysis
 - Safety Data Sheets (SDSs; also known as MSDSs)

Note: For SDSs for reagents and chemicals from other manufacturers, contact the manufacturer.

Limited product warranty

Life Technologies Corporation and/or its affiliate(s) warrant their products as set forth in the Life Technologies' General Terms and Conditions of Sale at www.thermofisher.com/us/en/home/global/terms-and-conditions.html. If you have any questions, please contact Life Technologies at www.thermofisher.com/support.

References

Kwok, S., and Higuchi, R. 1989. Avoiding false positives with PCR. *Nature* 339:237–238.

