

# **Brilliant III Ultra-Fast QRT-PCR Master Mix**

# **Quick Reference Guide for the ABI 7500 Fast Real-Time PCR System**

This quick reference guide provides an optimized protocol for using the Stratagene Brilliant III Ultra-Fast QRT-PCR Master Mix with the 7500 Fast Real-Time PCR System from Applied Biosystems. For detailed instructions, refer to the full product manual.

## Prepare the Reactions

- 1 Dilute the reference dye 1:500 using nuclease-free PCR-grade water.
- **2** Prepare the experimental reactions by combining the components of the reagent mixture in the order listed in the table below. Prepare a single reagent mixture for replicate reactions (plus *at least* one reaction volume excess) using multiples of each component. *Keep reagent mixture on ice*.

Reagent Mixture	
Nuclease-free PCR-grade water to bring final volume to 20 μl (including RNA	
10 μl of 2× QRT-PCR Master Mix	
x μl of experimental probe at optimized concentration (100–600 nM)	
x μl of upstream primer at optimized concentration (200–600 nM)	
x μl of downstream primer at optimized concentration (200–600 nM)	
0.3 μl of diluted reference dye	
0.2 µl of 100 mM DTT	
1 μl of RT/RNase Block	

- **3** Gently mix the reagent mixture without creating bubbles, then distribute the mixture to the experimental reaction tubes. *Keep the reactions on ice.*
- 4 Add x  $\mu$ l of experimental RNA to each reaction to bring the final reaction volume to 20  $\mu$ l. The table below lists a suggested quantity range for different RNA templates.

RNA	Quantity per reaction
Total RNA	0.1 pg — 100 ng
mRNA	0.1 pg — 1 ng

**5** Mix the reactions without creating bubbles, then centrifuge briefly.

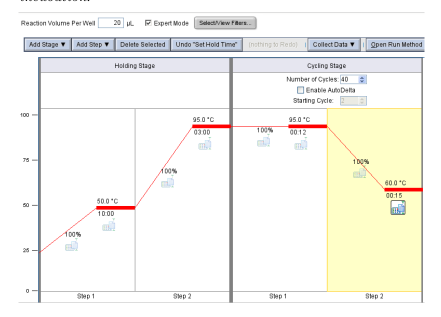


### Set Up the QPCR Plate and Thermal Profile

- 1 From the Home screen of the 7500 software, click Advanced Setup.
- 2 Complete the Setup screens for a new experiment as needed.

On the Experiment Properties screen, select **TaqMan Reagents** and the **Fast** ramp speed.

- **3** On the **Run Method** screen, set the reaction volume to 20 µl and mark the **Expert Mode** check box. Click **Select/View Filters** and deselect any filters not in use in the experiment.
- **4** Adjust the thermal profile according to the image below. *Note that a new step needs to be added to the beginning of the profile for the 50°C incubation.*



# Run the PCR

- 1 Place the reactions in the 7500 instrument.
- Program 2 Click START RUN.

**Analyze Data** 1 Analyze the results of the run as needed for your experiment.

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### **Product Information**

Catalog #600884, 400 reactions Catalog #600885, 4000 reactions

### **Ordering Information**

By phone (US only\*): 800-424-5444, x3 On the web: www.genomics.agilent.com

### **Technical Services**

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