

# Brilliant III Ultra-Fast SYBR<sup>®</sup> Green QPCR Master Mix

## Quick Reference Guide for the Stratagene Mx3000P/Mx3005P QPCR Systems

*This quick reference guide provides an optimized protocol for using the Stratagene Brilliant III Ultra-Fast SYBR<sup>®</sup> Green QPCR Master Mix with the Stratagene Mx3000P and Mx3005P QPCR Systems from Agilent. 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.

Reagent Mixture
Nuclease-free PCR-grade water to bring final volume to 20 $\mu$ l (including DNA)
10 $\mu$ l of 2 $\times$ SYBR Green QPCR Master Mix
x $\mu$ l of upstream primer at optimized concentration (200–500 nM)
x $\mu$ l of downstream primer at optimized concentration (200–500 nM)
0.3 $\mu$ l of diluted reference dye

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

DNA	Quantity per reaction
Genomic DNA	5 pg – 50 ng
cDNA	0.5 pg – 100 ng*

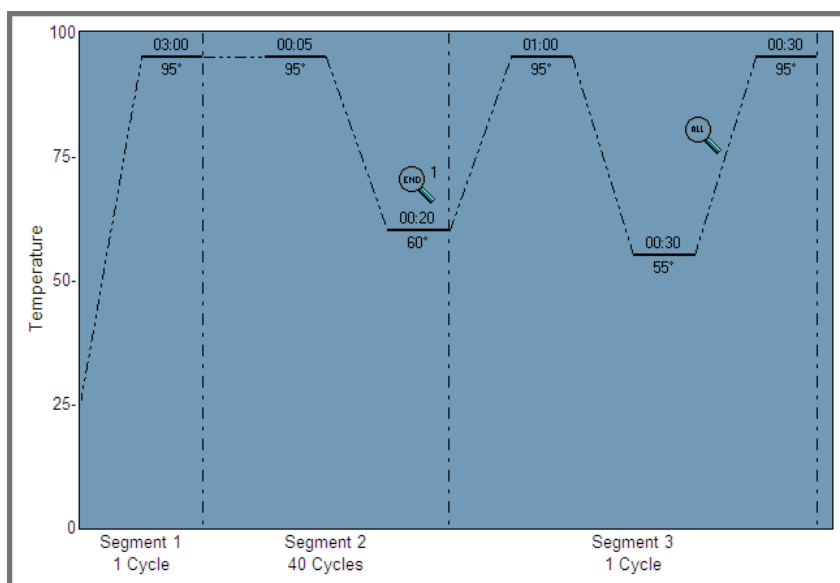
\*Refers to RNA input amount during cDNA synthesis

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



## Set Up the QPCR Plate and Thermal Profile

- 1 Complete the **Plate Setup** screen for a new experiment as needed, including assigning well types and assay information.
- 2 On the **Thermal Profile Setup** screen, set the **Thermal Profile Design** selection to **Standard**.
  - Under **Pre-Melt/RT Segment**, click **1 Plateau**.
  - Under **Amplification Segment**, click **Fast 2 Step**.
  - Under **Dissociation/Melt Segment**, click **Dissociation/Melt**.
- 3 Adjust the thermal profile according to the image below. The profile includes a 5-second denaturation step. Note that some assays may require a denaturation of up to 20 seconds. The exact denaturation time needs to be optimized for each target.



## Run the PCR Program

- 1 Place the reactions in the Mx3000P/Mx3005P instrument.
- 2 On the **Run** screen, click **Start Run**.

## Analyze Data

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

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### Product Information

Catalog #600882, 400 reactions  
Catalog #600883, 4000 reactions

### Ordering Information

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

### Technical Services

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