

EverGreen Universal qPCR Master Mix (2X)

Product Manual Version 1.2

This product is intended for research use only. Do not use this product for diagnostic purposes.



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Intended use

This product is intended for real-time qPCR applications using the intercalated dye chemistry for research purposes only. Please do not use this product in diagnostic applications.

We have optimized this product to be compatible with all commercially available realtime PCR machines.

Principle

Real-time qPCR is modification of the traditional PCR technique, in which amplification of DNA or cDNA (which is prepared from an RNA template) is monitored in real time by fluorescence.

Traditionally, the PCR products are run on a gel then visualized under a transilluminator. This type of analysis is termed end-point analysis because it only reflects the final reaction products. On the other hand, incorporation of a specific type of intercalating dyes in the reaction mix allows to monitor the reaction progression after each cycle. These intercalating dyes do not participate in the reaction, but bind strongly to double-stranded DNA. Upon binding, their fluorescence increases approximately 1,000 fold. Thus, after each replication, or doubling, of the targeted DNA element, the emitted fluorescence due to binding is also doubled. Fluorescence is then measured after each cycle by the real-time PCR machine, and the amount of the starting template can be inferred from fluorescence data.

This product contains all components needed for an efficient real-time qPCR reaction: a robust DNA polymerase, pure dNTPs, stabilizers, enhancers, and the intercalating dye. The user only needs to add the template and our optimized assays (or their own custom assays).

Kit contents and storage conditions

This kit contains 5 tubes of 1 mL EverGreen Universal qPCR Master Mix. Store at -20° C immediately upon receipt. Minimize exposure to light.



Running qPCR reactions

- 1. Let the EverGreen Universal qPCR Master Mix, Haven's optimized custom assays (or the user designed primers) and templates thaw completely at room temperature then put them on ice; for optimal results, work under minimal lighting and finish setting the experiment as quickly as possible
- 2. Determine the number of reactions needed for each specific assay

Example: if you wish to run triplicate reactions for 10 different samples, you will perform:

 $3_{replicates} X 10_{samples} = 30_{qPCR reactions}$

3. Prepare an assay master mix sufficient for the total number of reactions (determined in step2) plus 10% to account for pipetting errors; this is achieved by mixing the following components for final reaction volumes of 20 µL (<u>do not add the DNA/cDNA samples yet</u>):

Component	Volume per reaction	Example: Volume per 30+3 reactions	Final concentration
EverGreen qPCR MM	10 μL	330 μL	1 X
Haven's custom assay (20X)	lμL	33 μL	1 X
Template (DNA or cDNA)	Variable	Variable	10 pg – 100 ng (cDNA) 1-10 ng (DNA)
PCR-grade water	Up to 20 μL	Up to 660 μL	_

If user designed primers are used, refer to the following table:

Component	Volume per reaction	Example: Volume per 30+3 reactions	Final concentration
EverGreen qPCR MM	10 μL	330 μL	1 X
Forward primer	Variable	Variable	200 – 800 nM
Reverse primer	Variable	Variable	200 – 800 nM
Template (DNA or cDNA)	Variable	Variable	10 pg – 100 ng (cDNA) 1-10 ng (DNA)
PCR-grade water	Up to 20 μL	<i>Up to 660 μL</i>	_



- 4. If you are testing more than one assay, prepare an assay master mix for each assay as in step3
- 5. Load the appropriate amount of the assay master mix into the allocated PCR plate wells or PCR tubes

<u>Example:</u> if the amount of DNA/cDNA template was determined in step3 to be $2~\mu$ L, load $18~\mu$ L of the assay master mix into each allocated well or PCR tube

- Add the appropriate amount of DNA/cDNA template, as determined in step3, to each allocated well
- 7. Program the real-time thermal cycler as follows:

Step	Temperature	Time	Cycles
Initial denaturation	95°C	3 min	Hold
Denaturation	95°C	15 sec	40 cycles
Annealing/extension*	57-65°C**	60 sec	
Melting curve	65 to 95°C	0.2°C/sec ramp up rate	Hold

^{*} data collection in this step

8. Analyze data using your real-time thermal cycler's software

^{**} depending on the assay



Quality control

Each lot is tested for certain preset parameters to ensure compliance with manufacturing procedures.

Support

For questions, suggestions, or technical support, feel free to email our technical team at support@havensci.com.