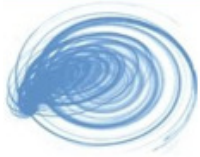


PrimerDesign Ltd

Custom designed real-time PCR assay with PerfectProbe™

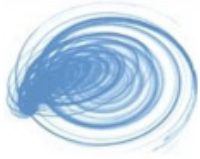
Instructions for detection and quantification of a target gene
using PerfectProbe™ detection chemistry





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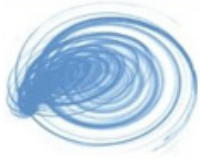
Introduction

This kit provides reagents for the target specific amplification of a target gene using real-time PCR with PerfectProbe™ detection chemistry. When resuspended, this kit provides primers that have been tested for priming specificity, and amplification efficiency at optimal concentrations. Real-time PCR is a very sensitive technology and it is not recommended to use more or less than the specified amount of primer and probe in each reaction. However, final reaction volumes between 15µl and 50µl are often successful and may be tested at the user's discretion. Unfortunately PrimerDesign is not able to provide technical support for protocols other than those provided.

For accurate gene expression measurements it is essential to normalise results from your quantitative real-time PCR experiments to a fixed reference. Normalising to a constitutively expressed housekeeping gene is the most common method. PrimerDesign provides researchers with a range of high quality methods and assays for housekeeping gene expression data.

- Normalising to a constitutively expressed housekeeping gene is the most common method. PrimerDesign provides researchers with a range of high quality real-time PCR assays for housekeeping gene expression data.
- geNorm™ is a system for selecting optimal housekeeping genes for any biological system (e.g. cell line, tissues sample)

Both can be found at www.primerdesign.co.uk



Kit Contents

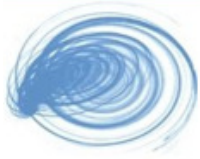
- **Lyophilised primer and probe set for a target gene selected by the user (BROWN)**
Includes primers and probe for **300, 600 or 900** 20µl reactions. The primers are individually designed and tested for priming specificity and amplification efficiency.
- **RNAse/DNAse free water (WHITE)**

Reagents and Equipment to Be Supplied by User

- **Real-Time PCR instrument**
- **Mastermix or Mastermix components**
This kit is designed to work well with all commercially available Mastermixes. However, we recommend the use of PrimerDesign 2x *Precision*[™] Mastermix.
- **Pipettors and Tips**
- **Vortex and centrifuge**
- **cDNA template**
The quality of cDNA will directly affect the quality of data generated using this kit. PrimerDesign recommends the use of the PrimerDesign *Precision*[™] RNA Reverse Transcription kit to generate cDNA from RNA.

Kit Storage

This PrimerDesign kit should be stored at -20°C on arrival. Freeze/thawing cycles should be kept to a minimum once resuspended. Under these conditions reagents are stable for six months from date of resuspension.



Licensing Agreement and Limitations of Use

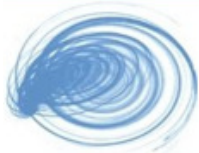
PCR is covered by several patents owned by Hoffman-Roche Inc and Hoffman-LaRoche, Ltd. Purchase of PrimerDesign kits does not include or provide licence with respect to any patents owned by Hoffman-La Roche or others.

PrimerDesign Satisfaction Guarantee

PrimerDesign takes pride in the quality of all of our products. Should this product fail to perform satisfactorily when used according to the protocols in this manual, PrimerDesign will replace the item free of charge.

Quality Control

As part of our routine quality assurance programme, all PrimerDesign products are monitored to ensure the highest levels of performance and reliability.



Bench-side Protocol

To minimise the risk of contamination with foreign DNA, we recommend that all pipetting be performed in a PCR clean environment. Ideally this would be a designated PCR cabinet. Filter tips are recommended for all pipetting steps.

1. Pulse-spin each tube in a centrifuge before opening.

This will ensure lyophilised primer and probe mix is in the base of the tube and is not spilt upon opening the tube.

2. Resuspend lyophilised primer and probe mix in RNase/DNase free water provided.

To ensure complete resuspension, vortex each tube thoroughly, allow to stand for 5 minutes and vortex again before use.

Reaction Number	Resuspension Volume
300 rxn	330 µl
600 rxn	660 µl
900 rxn	990 µl

There is a 10% over pipette in each kit

3. When using PrimerDesign 2X Precision™ Mastermix. Make up a mix containing all qPCR reagents according to the protocol below.

Component	1 Reaction
Resuspended primer/probe mix	1 µl*
PrimerDesign 2X Precision™ Mastermix	10 µl
RNase/DNase free water	4 µl
Final volume	15 µl

*working concentration of primers = 300nM in a 20µl reaction

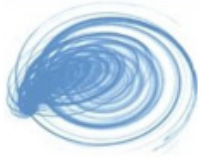
4. Pipette 15µl of this mix into each well according to your real-time PCR experimental plate set up.

5. Prepare cDNA for each of your samples (suggested concentration 5ng/µl) in RNase/DNase free water.

If the concentration of cDNA is not known, then dilute your RT reactions 1:10 (10µl of RT and 90µl of water)

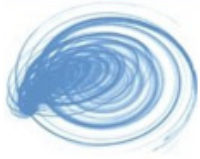
6. Pipette 5µl of diluted cDNA into each well, according to your experimental plate set up.

The final volume in each well is 20µl.



7. Include the following negative control wells.

- Include wells where the cDNA is replaced with RNase/DNase free water. Any amplification in this sample is indicative of cDNA cross contamination between wells, or contamination of one or more reagents.
- Include wells where the equivalent concentration of RNA is added minus the reverse transcription step. Amplification of these wells may indicate genomic DNA contamination of your RNA sample. A DNase treatment step is highly recommended during RNA extraction to prevent this occurring.



Amplification Protocol

Amplification conditions using **PrimerDesign 2X Precision™ MasterMix**.

Please note these are not standard Taqman® cycling conditions

	Step	Time	Temp
	Enzyme activation	10min	95°C
Cycling x50	Denaturation	15s	95°C
	DATA COLLECTION*	30s**	50°C
	Extension	15s	72°C

*Fluorogenic data should be collect during this step through the FAM channel.

** For some real-time PCR platforms 15 seconds is sufficient to acquire data through 2 channels. This step may be shortened accordingly. A 30s step is recommended for the Bio-Rad iCycler.