

# GeneQuery<sup>TM</sup> Human Apoptosis qPCR Array Kit (GQH-APO)

Catalog #GK001

#### **Product Description**

ScienCell's GeneQuery<sup>TM</sup> Human Apoptosis qPCR array kit (GQH-APO) is designed to facilitate gene expression profiling of key genes involved in human apoptosis. 88 genes are selected in this kit based on database and literature research.

GeneQuery<sup>TM</sup> qPCR array kits are qPCR ready in a 96-well plate format, with each well containing one primer set that can specifically recognize and efficiently amplify a target gene's cDNA. The carefully designed primers ensure that: (i) the optimal annealing temperature in qPCR analysis is 65°C (with 2 mM Mg<sup>2+</sup>, and no DMSO); (ii) the primer set recognizes all known transcript variants of target gene, unless otherwise indicated; and (iii) only one gene is amplified. Each primer set has been validated by qPCR with melt curve analysis, and gel electrophoresis.

## GeneQuery<sup>TM</sup> qPCR Array Kit Controls

Each GeneQuery<sup>TM</sup> plate contains eight controls (Figure 1).

- Five target housekeeping genes (β-actin, GAPDH, LDHA, NONO, and PPIH), which enable normalization of data.
- The Genomic DNA (gDNA) Control (GDC) detects possible gDNA contamination in the cDNA samples. It contains a primer set targeting a non-transcribed region of the genome.
- Positive PCR Control (PPC) tests whether samples contain inhibitors or other factors that may negatively affect gene expression results. The PPC consists of a predispensed synthetic DNA template and a primer set that can amplify it. The sequence of the DNA template is not present in the human genome, and thus tests the efficiency of the polymerase chain reaction itself.
- The No Template Control (NTC) is strongly recommended, and can be used to monitor the DNA contamination introduced during the workflow such as reagents, tips, and the lab bench.

#### **Kit Components**

Component	Quantity	Storage
GeneQuery <sup>TM</sup> array plate with lyophilized primers	1	4°C or -20°C
Optical PCR plate seal	1	RT
Nuclease-free H <sub>2</sub> O	2 mL	4°C

#### Additional Materials Required (Materials Not Included in Kit)

Component	Recommended
Reverse transcriptase	MultiScribe Reverse Transcriptase (Life Tech, Cat. #4311235)
cDNA template	Customers' samples
qPCR master mix	FastStart Essential DNA Green Master (Roche, Cat. #06402712001)

## **Quality Control**

All the primer sets are validated by qPCR with melt curve analysis. The PCR products are analyzed by gel electrophoresis. Single band amplification is confirmed for each set of primers.

#### **Product Use**

GQH-APO is for research use only. It is not approved for human or animal use, or for application in clinical or *in vitro* diagnostic procedures.

## **Shipping and Storage**

The product is shipped at ambient temperature. Upon receipt, the plate should be stored at 4°C and is good for up to 12 months. For long-term storage (>1 year), store the plate at -20°C in a manual defrost freezer.

**Note:** The primers in each well are lyophilized.

- 1. Prior to use, allow plates to warm to room temperature.
- 2. Briefly centrifuge at 1,500x g for 1 minute before slowly peeling off the seal.
- 3. Prepare 20 µl PCR reactions for one well as shown in Table 1.

Table 1

cDNA template	0.2 – 250 ng
2x qPCR master mix	10 μ1
Nuclease-free H <sub>2</sub> O	variable
Total volume	20 μl

*Important: Only* use polymerases with hot-start capability to prevent possible primer-dimer formation. *Only* use nuclease-free reagents in PCR amplification.

4. Add the mixture of 2x qPCR master mix, cDNA template, and nuclease-free H<sub>2</sub>O to each well containing the lyophilized primers. Seal the plate with the provided optical PCR plate seal.

Important: In NTC control well, do NOT add cDNA template. Add 2x qPCR master mix and nuclease-free H2O only.

- 5. Briefly centrifuge the plates at 1,500x g for 1 minute at room temperature. For maximum reliability, replicates are strongly recommended (minimum of 3).
- 6. For PCR program setup, please refer to the instructions of the master mix of the user's choice. We recommend a typical 3-step qPCR protocol for a 200nt amplicon:

Three-step cycling protocol

Step	Temperature	Time	Number of cycles
Initial denaturation	95°C	10 min	1
Denaturation	95°C	20 sec	
Annealing	65°C	20 sec	40
Extension	72°C	20 sec	40
Data acquisition	Plate read		
Recommended	Melting cı	ırve analysis	1
Hold	4°C	Indefinite	1

7. (Optional) Load the PCR products on 1.5% agarose gel and perform electrophoresis to confirm the single band amplification in each well.

Figure 1. Layout of GeneQuery<sup>TM</sup> qPCR array kit controls.

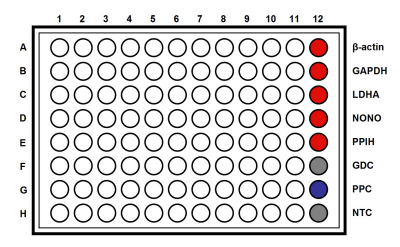


Table 2. Interpretation of control results:

Controls	Results	Interpretation	Suggestions
Housekeeping gene controls	Variability of a housekeeping gene's Cq value	The expression of the housekeeping gene is variable in samples; cycling program is incorrect	Choose a constantly expressed target, or analyze expression levels of multiple housekeeping genes; use correct cycling program and make sure that all cycle parameters have been correctly entered
gDNA Control (GDC)	Cq ≥ 35	No gDNA detected	N/A
	Cq < 35	The sample is contaminated with gDNA	Perform DNase digestion during RNA purification step
Positive PCR Control (PPC)	Cq > 30; or The Cq variations > 2 between qPCR Arrays.	Poor PCR performance; possible PCR inhibitor in reactions; cycling program incorrect	Eliminate inhibitor by purifying samples; use correct cycling program and make sure that all cycle parameters have been correctly entered
No Template Control (NTC)	Positive	DNA contamination in workflow	Eliminate sources of DNA contamination (reagents, plastics, etc.)

Figure 2. A typical amplification curve showing the amplification of a qPCR product.

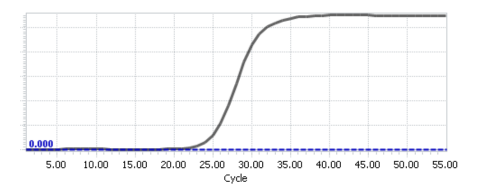
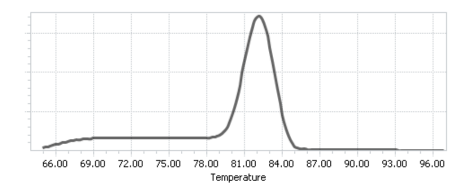


Figure 3. A typical melting peak of a qPCR product.



### Quantification Method: Comparative $\Delta\Delta$ Cq (Quantification Cycle Value) Method

1. **Note:** Please refer to your qPCR instrument's data analysis software for data analysis. The method provided here serves as guidance for quick manual calculations.

You can use one or more housekeeping genes as a reference to normalize samples.

*Important:* We highly recommend using all 5 housekeeping genes included in this kit,  $\beta$ -actin, GAPDH, LDHA, NONO, and PPIH.

2. For a single housekeeping gene,  $\Delta$ Cq (ref) is the quantification cycle number change for that housekeeping gene (HKG) between an experimental sample and control sample.

$$\Delta$$
Cq (ref) = Cq (HKG, experimental sample) - Cq (HKG, control sample)

When using multiple housekeeping genes as a reference, we recommend normalizing using the geometric mean [1] of the expression level change, which is the same as normalizing using the arithmetic mean of  $\Delta Cq$  of the selected housekeeping genes.

 $\Delta$ Cq (ref) = average ( $\Delta$ Cq (HKG1),  $\Delta$ Cq (HKG2),.....,  $\Delta$ Cq (HKG n)) (n is the number of housekeeping genes selected)

*If* using all 5 housekeeping genes included in this kit, β-actin, GAPDH, LDHA, NONO, and PPIH, use the following formula:

$$\Delta$$
Cq (ref) = ( $\Delta$ Cq( $\beta$ -actin)+ $\Delta$ Cq(GAPDH)+ $\Delta$ Cq(LDHA)+ $\Delta$ Cq(NONO)+ $\Delta$ Cq(PPIH)) /5

*Note:*  $\Delta$ Cq (HKG) = Cq (HKG, experimental sample) - Cq (HKG, control sample), and  $\Delta$ Cq (HKG) value can be positive, 0, or negative.

3. For any of your genes of interest (GOI),

$$\Delta$$
Cq (GOI) = Cq (GOI, experimental sample) - Cq (GOI, control sample)

$$\Delta\Delta$$
Cq =  $\Delta$ Cq (GOI) -  $\Delta$ Cq (ref)

Normalized GOI expression level fold change =  $2^{-\Delta\Delta Cq}$ 

#### References

[1] Vandesompele J, De Preter K, Pattyn F, Poppe B, Van Roy N, De Paepe A, Speleman F. (2002) "Accurate normalization of real-time quantitative RT-PCR data by geometric averaging of multiple internal control genes." *Genome Biol.* 3(7): 1-12.

### Example: Comparative ΔΔCq (Quantification Cycle Value) Method

Table 3. Cq (Quantification Cycle) values of 2 genes-of-interest and 5 housekeeping genes obtained for experimental and control samples.

	Genes o	f Interest		House	keeping G	enes	
Samples	GOI1	GOI2	β-actin	GAPDH	LDHA	NONO	PPIH
Experimental	21.61	22.19	17.16	17.84	20.12	19.64	26.40
Control	33.13	26.47	18.20	18.48	20.57	19.50	26.55

$$\Delta$$
Cq (ref) = ( $\Delta$ Cq( $\beta$ -actin)+ $\Delta$ Cq(GAPDH)+ $\Delta$ Cq(LDHA)+ $\Delta$ Cq(NONO)+ $\Delta$ Cq(PPIH)) /5 = ((17.16-18.20)+(17.84-18.48)+(20.12-20.57)+(19.64-19.50)+(26.40-26.55))/5 = -0.43

$$\Delta$$
Cq (GOI1) = 21.61 - 33.13  
= -11.52

$$\Delta$$
Cq (GOI2) = 22.19 - 26.47  
= -4.28

$$\Delta\Delta$$
Cq (GOI1) =  $\Delta$ Cq (GOI1) -  $\Delta$ Cq (ref)  
= -11.52 - (-0.43)  
= -11.09

$$\Delta\Delta$$
Cq (GOI2) =  $\Delta$ Cq (GOI2) -  $\Delta$ Cq (ref)  
= -4.28 - (-0.43)  
= -3.85

Normalized GOI1 expression level fold change = 
$$2^{-\Delta\Delta Cq \text{ (GOI1)}}$$
  
=  $2^{11.09}$   
= 2180

Normalized GOI2 expression level fold change = 
$$2^{-\Delta\Delta Cq \text{ (GOI2)}}$$
  
=  $2^{3.85}$   
= 14.4

Conclusion: Upon treatment, expression level of GOI1 increased 2,180 fold, and expression level of GOI2 increased 14.4 fold.



# GeneQuery<sup>TM</sup> Human Apoptosis qPCR Array Kit (GQH-APO)

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GeneQuery<sup>TM</sup> Human Apoptosis qPCR Array Plate Layout\* (8 controls in Bold and Italic)

	1	2	3	4	5	6	7	8	9	10	11	12
A	ABL1	BAG1	BIRC1	BRAF	CASP6	DFFA	IL1RAP	MYD88	PIK3R3	RELA	TNFRSF9	β-actin
В	AIFM1	BAG3	BIRC2	CAPN1	CASP9	DIABLO	IL3	NGF	PPP3CA	RIPK1	TNFSF10	GAPDH
C	AKT1	BCL10	BIRC3	CAPN2	CD40	ENDOG	IRAK2	NOD1	PPP3CB	RIPK2	TP53	LDHA
D	AKT2	BCL2	BIRC5	CASP1	CFLAR	FAS	IRAK3	NOL3	PPP3R1	TNF	TP53BP2	NONO
$\mathbf{E}$	AKT3	BCL2L1	BIRC6	CASP10	CHUK	HRK	IRAK4	PIK3CB	PRKAR1A	TNFRSF10B	TRADD	PPIH
$\mathbf{F}$	APAF1	BCL2L10	BNIP2	CASP2	CIDEA	IGF1R	LTBR	PIK3CG	PRKAR2B	TNFRSF10D	TRAF2	GDC
G	ATM	BID	BNIP3	CASP3	CRADD	IL1B	MAP3K14	PIK3R1	PRKX	TNFRSF11B	TRAF3	PPC
H	BAD	BIK	BNIP3L	CASP4	DAPK1	IL1R1	MCL1	PIK3R2	PYCARD	TNFRSF1B	XIAP	NTC

<sup>\*</sup> gene selection may be updated based on new research and development

# Plate type A

Brand	Model	kit catalog #
ABI / Life Tech	ABI 5700	GK001-A
	ABI 7000	GK001-A
	ABI 7300	GK001-A
	ABI 7500	GK001-A
	ABI 7700	GK001-A
	ABI 7900 HT	GK001-A
	QuantStudio	GK001-A
	ViiA 7	GK001-A
Bio-Rad	Chromo4	GK001-A
	iCycler	GK001-A
	iQ5	GK001-A
	MyiQ	GK001-A
	MyiQ2	GK001-A
Eppendorf / Life Tech	Matercycler ep realplex 2	GK001-A
	Matercycler ep realplex 4	GK001-A
Stratagene	MX3000P	GK001-A
J	MX3005P	GK001-A

## Plate type B

Brand	Model	kit catalog #
ABI / Life Tech	ABI 7500 Fast	GK001-B
	ABI 7900 HT Fast	GK001-B
	QuantStudio Fast	GK001-B
	StepOnePlus	GK001-B
	ViiA 7 Fast	GK001-B
	27/2	01/001 5
Bio-Rad	CFX Connect	GK001-B
	CFX96	GK001-B
	DNA Engine Opticon 2	GK001-B
Stratagene	MX4000	GK001-B

# Plate type C

Brand	Model	kit catalog #
Roche	Lightcycler 96	GK001-C
	Lightcycler 480 (96-well)	GK001-C