

## TECHNICAL DATASHEET

PRODUCT NAME

Mouse Hhex promoter primer Pairs

Official full name: Hematopoietically expressed homeobox

Other name: Hex, Prh, Prhx, Hhex-rs2

Primary source: MGI: 96086

 Cat. No: pp-1027-050
 Size: 50 μl
 Concentration: 10 μM
 Lot #: 001

 Cat. No: pp-1027-500
 Size: 500 μl
 Concentration: 10 μM
 Lot #: 001

**10 sets of our primer pairs:** 50 μl (see our list)

500 μl

**Description:** The primer pair cat:# pp-1027 (-050, -500) is specific to a DNA region in the mouse Hhex gene promoter [1]. These primers can be used to amplify DNA isolated by chromatin immunoprecipitation (ChIP). Primers are optimized to be used in quantitative polymerase chain reaction (qPCR) (**Figures 1, 2 and 3**). See overview below.

Specificity: Mouse: positive

Other species: not tested

Expected PCR product size: 125 base pairs (bp).

Format: In solution in MiliQ water at the concentration of 10  $\mu M$  (each oligonucleotide primer is at the final

concentration of 5  $\mu$ M).

**Storage:** For long storage, store at -20°C. Avoid multiple freeze-thaw cycles.

Precautions: This product is for research use only. Not for use in diagnostic or therapeutic procedures.

References: [1] O'Neill L.P., VerMilyea M.D. and Turner B.M. (2006) Nat. Genet. 38(7):835-41.

[2] Guo Y., Chan R, Ramsey H., Li W., Xie X., Shelley W.C., Martinez-Barbera J.P., Bort B., Zaret K., Yoder M. and Hromas R. (2003) Blood 102 (7):2428-35

Last data sheet update: September 14, 2007

Availability date: September 03, 2007

Lot #: 001/ day of the synthesis: May 25, 2007/ day of QC: August 17, 2007/ aliquoting day: August 24, 2007

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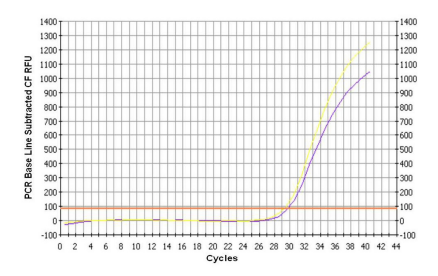


Figure 1

DNA from mouse fibroblast 3T3 cells was analyzed in duplicate by real-time PCR starting from 5  $\mu$ l of DNA template (0.03  $\mu$ g/ml) using the Diagenode primers to amplify a region in the mouse Hhex gene promoter (cat#: pp-1027-050, -500). One  $\mu$ l of provided primer pairs is used by PCR of 25  $\mu$ l final volume. A Real-Time PCR Detection System and iQ SYBR Green have been used. qPCR conditions used are as follows: 95°C for 3 minutes, 41 cycles of: [95°C for 60 seconds, 60°C for 60 seconds and 72°C for 90 seconds]. Duplicates are shown in yellow and purple. Threshold position is in orange.

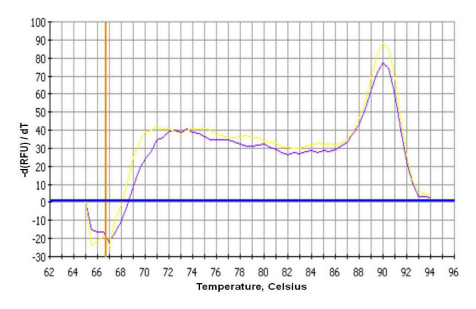


Figure 2.

Melting curves obtained with the primers cat#: pp1027 (-050, -500) used in the above qPCR. Conditions were 60 cycles of 65°C for 1 minute and increment of 0.5°C per cycle. Duplicates are shown in yellow and purple.



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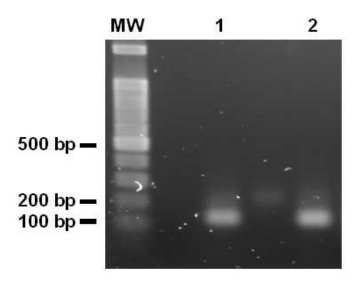


Figure 3

qPCR products were analysed by electrophoresis (1.5% agarose gel) stained with SYBR Safe and illuminated with UV light. The left lane shows molecular weight markers (MW) that decrease in size by 100 bp. Different qPCR products using different primer pairs which are available at Diagenode were tested: 1: primers for mouse Cdx2 gene promoter (pp-1025-050, -500), 2: primers for mouse Hhex gene promoter (pp-1027-050, -500). For more details about other primer pairs, see data sheet.

**Overview:** In the adult, the Hhex gene is preferentially expressed in hematopoietic cells but it is generally down-regulated during terminal differentiation of hematopoietic cells. The lack of Hhex is detrimental to the differentiation of the hemangioblast to hematopoietic progenitors and to a lesser extent, to endothelial cells [2].