

## **TGIF Polyclonal Antibody**

Catalog No: YT4635

**Reactivity:** Human; Mouse; Rat

**Applications:** WB;ELISA

Target: TGIF

**Fields:** >>TGF-beta signaling pathway

Q15583

P70284

Gene Name: TGIF1

Protein Name: Homeobox protein TGIF1

Human Gene Id: 7050

**Human Swiss Prot** 

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No:

Mouse Gene ld: 21815

**Mouse Swiss Prot** 

No:

**Immunogen:** Synthesized peptide derived from the C-terminal region of human TGIF.

**Specificity:** TGIF Polyclonal Antibody detects endogenous levels of TGIF protein.

**Formulation :** Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

Source: Polyclonal, Rabbit, IgG

**Dilution:** WB 1:500 - 1:2000. ELISA: 1:10000. Not yet tested in other applications.

**Purification:** The antibody was affinity-purified from rabbit antiserum by affinity-

chromatography using epitope-specific immunogen.

Concentration: 1 mg/ml

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Storage Stability: -15°C to -25°C/1 year(Do not lower than -25°C)

Observed Band: 43kD

**Background :** The protein encoded by this gene is a member of the three-amino acid loop

extension (TALE) superclass of atypical homeodomains. TALE homeobox proteins are highly conserved transcription regulators. This particular homeodomain binds to a previously characterized retinoid X receptor responsive element from the cellular retinol-binding protein II promoter. In addition to its role in inhibiting 9-cis-retinoic acid-dependent RXR alpha transcription activation of the retinoic acid responsive element, the protein is an active transcriptional corepressor of SMAD2 and may participate in the transmission of nuclear signals during development and in the adult. Mutations in this gene are associated with holoprosencephaly type 4, which is a structural anomaly of the brain. Alternative splicing has been observed at this locus and multiple splice variants encoding

distinct isoforms are described. [provide

**Function:** disease:Defects in TGIF1 are the cause of holoprosencephaly type 4 (HPE4)

[MIM:142946]. Holoprosencephaly (HPE) [MIM:236100] is the most common structural anomaly of the brain, in which the developing forebrain fails to correctly separate into right and left hemispheres. Holoprosencephaly is genetically heterogeneous and associated with several distinct facies and phenotypic variability.,function:Binds to a retinoid X receptor (RXR) responsive element from the cellular retinol-binding protein II promoter (CRBPII-RXRE). Inhibits the 9-cisretinoic acid-dependent RXR alpha transcription activation of the retinoic acid responsive element. Active transcriptional corepressor of SMAD2. Links the nodal signaling pathway to the bifurcation of the forebrain and the establishment of ventral midline structures. May participate in the transmission of nuclear signals during development and in the adu

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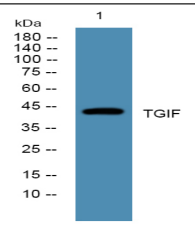
Nucleus.

Subcellular Location :

**Expression :** Brain, Liver, Placenta,

## **Products Images**

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Western blot analysis of lysates from KB cells, primary antibody was diluted at 1:1000, 4° over night