

Thyroid transcription factor-1 (TTF1) (PT0101R) rabbit mAb

Catalog No: YM7238

Reactivity: Human; Mouse; (predicted: Rat)

Applications: IHC;WB; ELISA

Target: TTF-1

Gene Name: NKX2-1

Protein Name: AV026640;BCH;Benign chorea;BHC;Homeobox protein NK 2 homolog

A;Homeobox protein NK-2 homolog A;Homeobox protein Nkx 2.1;Homeobox protein Nkx-2.1;Homeobox protein Nkx2.1;NK 2;NK 2 homolog A;NK2;NK2 hom

Human Swiss Prot

No:

Mouse Swiss Prot

No:

Rat Swiss Prot No: P23441

Immunogen: Synthesized peptide derived from human TTF1 AA range:50-150

Specificity: This antibody detects endogenous levels of TTF-1

Formulation: PBS, 50% glycerol, 0.05% Proclin 300, 0.05%BSA

Source: Monoclonal, Rabbit IgG1, Kappa

P43699

P50220

Dilution: IHC 1:100-500, WB 1:500-1000, ELISA 1:5000-20000

Purification: Recombinant Expression and Affinity purified

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

Molecularweight: 41kD

Background : This gene encodes a protein initially identified as a thyroid-specific transcription

factor. The encoded protein binds to the thyroglobulin promoter and regulates the



expression of thyroid-specific genes but has also been shown to regulate the expression of genes involved in morphogenesis. Mutations and deletions in this gene are associated with benign hereditary chorea, choreoathetosis, congenital hypothyroidism, and neonatal respiratory distress, and may be associated with thyroid cancer. Multiple transcript variants encoding different isoforms have been found for this gene. This gene shares the symbol/alias 'TTF1' with another gene, transcription termination factor 1, which plays a role in ribosomal gene transcription. [provided by RefSeq, Feb 2014],

Function:

disease:Defects in NKX2-1 are the cause of benign hereditary chorea (BHC) [MIM:118700]; also known as hereditary chorea without dementia. BHC is an autosomal dominant movement disorder. The early onset of symptoms (usully before the age of 5) and the observation that in some BHC families the symptoms tend to decrease in adulthood suggests that the disorder results from a developmental disturbance of the brain. BHC is non-progressive and patients have normal or slightly below normal intelligence. There is considerable inter- and intrafamilial variability, including dysarthria, axial distonia and gait disturbances., disease:Defects in NKX2-1 are the cause of choreoathetosis, hypothyroidism, and neonatal respiratory distress (CHNRD) [MIM:610978]. This syndrome include neurological, thyroid, and respiratory problems., function:Transcription factor that binds and activates the promoter of thyro

Subcellular Location:

Nuclear

Expression: Thyroid/ Lung

Tag: recombinant

Sort: 999

No4: 1

Host: Rabbit

Modifications: Unmodified

Products Images

2/2