

## TRβ1 (phospho Ser142) Polyclonal Antibody

Catalog No :	YP0479		
Reactivity :	Human;Mouse;Rat		
Applications :	WB;ELISA		
Target :	ΤRβ1		
Fields :	>>Neuroactive ligand-receptor interaction;>>Thyroid hormone signaling pathway		
Gene Name :	THRB		
Protein Name :	Thyroid hormone receptor beta		
Human Gene Id :	7068		
Human Swiss Prot No :	P10828		
Mouse Gene Id :	21834		
Mouse Swiss Prot No :	P37242		
Rat Swiss Prot No :	P18113		
Immunogen :	The antiserum was produced against synthesized peptide derived from human TR-beta1 around the phosphorylation site of Ser142. AA range:116-165		
Specificity :	Phospho-TR $\beta$ 1 (S142) Polyclonal Antibody detects endogenous levels of TR $\beta$ 1 protein only when phosphorylated at S142.		
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: 1:5000. 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		
Storage Stability :	-15°C to -25°C/1 year(Do not lower than -25°C)		
Storage Stability .			
Observed Band :	45kD		
Cell Pathway :	Neuroactive ligand-receptor interaction;		
Background :	The protein encoded by this gene is a nuclear hormone receptor for triiodothyronine. It is one of the several receptors for thyroid hormone, and has been shown to mediate the biological activities of thyroid hormone. Knockout studies in mice suggest that the different receptors, while having certain extent of redundancy, may mediate different functions of thyroid hormone. Mutations in this gene are known to be a cause of generalized thyroid hormone resistance (GTHR), a syndrome characterized by goiter and high levels of circulating thyroid hormone (T3-T4), with normal or slightly elevated thyroid stimulating hormone (TSH). Several alternatively spliced transcript variants encoding the same protein have been observed for this gene. [provided by RefSeq, Jul 2008],		
Function :	disease:Defects in THRB are the cause of generalized thyroid hormone resistance (GTHR) [MIM:188570, 274300]. GTHR is transmitted as an autosoma dominant trait, but an autosomal recessive form also exists. The disease is characterized by goiter, abnormal mental functions, increased susceptibility to infections, abnormal growth and bone maturation, tachycardia and deafness. Affected individuals may also have attention deficit-hyperactivity disorders (ADHD) and language difficulties. GTHR patients also have high levels of circulating thyroid hormones (T3-T4), with normal or slightly elevated thyroid stimulating hormone (TSH).,disease:Defects in THRB are the cause of selective pituitary thyroid hormone resistance (PRTH) [MIM:145650]; also called familial hyperthyroidism due to inappropriate thyrotropin secretion. PRTH is a variant form of thyroid hormone resistance and is characterized by c		
Subcellular Location :	Nucleus.		
Expression :	Brain,Kidney,Pituitary,Placenta,Testis,		
Tag:	orthogonal		
Sort :	23634		
No4 :	1		



Products Images				
TR-beta1 (pSer142)	293 293 117 85 48 34 26 19 (kD)	Western blot analysis of lysates from 293 cells treated with PMA 125ng/ml 30', using TR-beta1 (Phospho-Ser142) Antibody. The lane on the right is blocked with the phospho peptide.		