

BBS2 rabbit pAb

Catalog No: YT6543

Reactivity: Human; Mouse; Rat

Applications: WB

Target: BBS2

Gene Name: BBS2

Protein Name: BBS2

Human Gene Id: 583

Human Swiss Prot

No:

Mouse Gene ld: 67378

Q9BXC9

Q9CWF6

Mouse Swiss Prot

No:

Rat Gene Id: 113948

Rat Swiss Prot No: Q99MH9

Immunogen: Synthesized peptide derived from human BBS2 AA range: 149-199

Specificity: This antibody detects endogenous levels of BBS2 at Human/Mouse/Rat

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

Source: Polyclonal, Rabbit, IgG

Dilution: WB 1 ? 500-2000

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)

Molecularweight: 79kD

Background: This gene is a member of the Bardet-Biedl syndrome (BBS) gene family. Bardet-

Biedl syndrome is an autosomal recessive disorder characterized by severe pigmentary retinopathy, obesity, polydactyly, renal malformation and mental retardation. The proteins encoded by BBS gene family members are structurally diverse and the similar phenotypes exhibited by mutations in BBS gene family members is likely due to their shared roles in cilia formation and function. Many BBS proteins localize to the basal bodies, ciliary axonemes, and pericentriolar regions of cells. BBS proteins may also be involved in intracellular trafficking via microtubule-related transport. The protein encoded by this gene forms a multiprotein BBSome complex with seven other BBS proteins.[provided by

RefSeq, Oct 2014],

Function: disease:Defects in BBS2 are the cause of Bardet-Biedl syndrome type 2 (BBS2)

[MIM:209900]. Bardet-Biedl syndrome (BBS) is a genetically heterogeneous, autosomal recessive disorder characterized by usually severe pigmentary retinopathy, early onset obesity, polydactyly, hypogenitalism, renal malformation

and mental retardation. Secondary features include diabetes mellitus,

hypertension and congenital heart disease. A relatively high incidence of BBS is found in the mixed Arab populations of Kuwait and in Bedouin tribes throughout the Middle East, most likely due to the high rate of consaguinity in these populations and a founder effect..function:The BBSome complex is required for

ciliogenesis but is dispensable for centriolar satellite function. This ciliogenic function is mediated in part by the Rab8 GDP/GTP exchange factor, which

localizes to the basal body and contacts the BBSome. Rab8

Subcellular

Cell projection, cilium membrane. Cytoplasm. Cytoplasm, cytoskeleton, microtubule organizing center, centrosome, centriolar satellite.

Eocation:

Expression : Widely expressed.

Sort: 2599

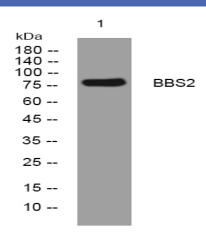
No4: 1

Host: Rabbit

Modifications: Unmodified



Products Images



Western blot analysis of lysates from DU145 cells, primary antibody was diluted at 1:1000, 4° over night