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BTR1 Rabbit pAb

CatalogNo: YT0545

Key Features

Host Species

Rabbit

Reactivity

Human,Mouse

ApplicationsWB,ELISA

MW • 100kD (Observed) lsotype • lgG

Recommended Dilution Ratios

WB 1:500-1:2000 ELISA 1:20000 Not yet tested in other applications.

Storage

Storage*-15°C to -25°C/1 year(Do not lower than -25°C)FormulationLiquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

Basic Information

Clonality Polyclonal

Immunogen Information

ImmunogenThe antiserum was produced against synthesized peptide derived from human SLC4A11.
AA range:291-340

Specificity BTR1 Polyclonal Antibody detects endogenous levels of BTR1 protein.

Target Information

Gene name SLC4A11

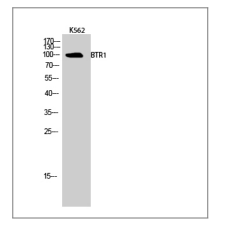
Protein Name Sodium bicarbonate transporter-like protein 11

| Organism | Gene ID | UniProt ID |
|----------|---------------|----------------|
| Human | <u>83959;</u> | <u>Q8NBS3;</u> |
| Mouse | | A2AJN7; |

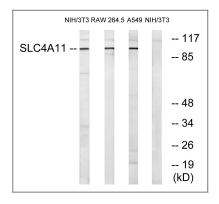
CellularCell membrane ; Multi-pass membrane protein . Basolateral cell membrane ; Multi-passLocalizationmembrane protein .

- **Tissue specificity** Widely expressed. Highly expressed in kidney, testis, salivary gland, thyroid, trachea and corneal endothelium. Not detected in retina and lymphocytes. ; [Isoform 3]: Expressed in corneal endothelium (at protein level). ; [Isoform 5]: The predominant isoform in corneal endothelium (at protein level).
- FunctionDisease:Defects in SLC4A11 are the cause of corneal dystrophy and perceptive deafness
(CDPD) [MIM:217400]; also known as corneal dystrophy and sensorineural deafness or
Harboyan syndrome. CDPD consists of congenital corneal endothelial dystrophy and
progressive perceptive deafness. Inheritance is autosomal recessive.,Disease:Defects in
SLC4A11 are the cause of corneal endothelial dystrophy type 2 (CHED2) [MIM:217700]; also
known as congenital hereditary endothelial dystrophy of cornea. This bilateral corneal
dystrophy is characterized by corneal opacification and nystagmus. Inheritance is
autosomal recessive.,Function:Transporter involved in borate homeostasis. In the absence
of borate, it functions as a Na(+) and OH(-)(H(+)) channel. In the presence of borate
functions as an electrogenic Na(+) coupled borate
cotransporter.,PTM:Glycosylated.,similarity:Belongs to the anion exchanger (TC 2.A.31)
family.,tissue specificity:Widely expressed. Highly expressed in kidney, testis, salivary
gland, thyroid, trachea and corneal endothelium. Not detected in retina and lymphocytes.,

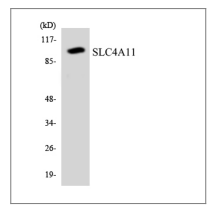
Validation Data



Western Blot analysis of K562 cells using BTR1 Polyclonal Antibody



Western blot analysis of lysates from NIH/3T3, RAW264.7, and A549 cells, using SLC4A11 Antibody. The lane on the right is blocked with the synthesized peptide.



Western blot analysis of the lysates from HT-29 cells using SLC4A11 antibody.

Contact information

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|------------|--|
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Please scan the QR code to access additional product information: **BTR1 Rabbit pAb**

For Research Use Only. Not for Use in Diagnostic Procedures.

Antibody | ELISA Kits | Protein | Reagents