bs-10314R

[Primary Antibody]

VNN1 Rabbit pAb



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- DATASHEFT				2 0000
Host: Ra	bbit Iso	type: IgG	Applications:	WB (1:500-2000)
Clonality: Po	lyclonal			IHC-P (1:100-500)
GenelD: 88	76 SI	VISS: 095497		IF (1:100-500)
Target: VN	N1			Flow-Cyt (2µg/Tes
Immunogen: KLH conjugated synthetic peptide derived from human VNN1: 151-250/513.		Reactivity:	Human, Mouse, Rat (predicted: Rabbit,	
Purification: aff	inity purified by Protein A			Cow)
Concentration: 1mg/ml			Predicted	
Storage: 0.(01M TBS (pH7.4) with 1% BSA, 0.	02% Proclin300 and 50%	MW.: ^{52 kDa}	
Sh	ipped at 4°C. Store at -20°C for o eze/thaw cycles.	one year. Avoid repeated	Subcellular Location:	Cell membrane
Background: Th sh bio as pa ha th ox lik ex mi clo tw Re	is gene encodes a member of th are extensive sequence similarit otinidase. The family includes se sociated proteins, a few of which rticipate in hematopoietic cell t s been demonstrated for any of ey possess pantetheinase activit idative-stress response. This pro- ely a GPI-anchored cell surface of pressed by the perivascular thy gration of T-cell progenitors to the soe proximity to, and in the same o other vanin genes on chromos fSeq, Feb 2009]	e vanin family of proteins, which cy with each other, and also with creted and membrane- n have been reported to rafficking. No biotinidase activity the vanin proteins, however, cy, which may play a role in otein, like its mouse homolog, is molecule. The mouse protein is nic stromal cells and regulates the thymus. This gene lies in e transcriptional orientation as, some 6q23-q24. [provided by		
- VALIDATION	IMAGES			

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Sample: placenta(Mouse) Lysate at 40 ug Primary: Anti-VNN1(bs-10314R) at 1/300 dilution Secondary: IRDye800CW Goat Anti-Rabbit IgG at 1/20000 dilution Predicted band size: 52 kD Observed band size: 52 kD



Tissue/cell: rat spleen tissue; 4% Paraformaldehyde-fixed and paraffinembedded; Antigen retrieval: citrate buffer ($0.01 \text{M}, \text{pH}\,6.0$), Boiling bathing for 15min; Block endogenous peroxidase by 3% Hydrogen peroxide for 30min; Blocking buffer (normal goat serum,C-0005) at 37°C for 20 min; Incubation: Anti-VNN1 Polyclonal Antibody, Unconjugated(bs-10314R) 1:200, overnight at 4°C, followed by conjugation to the secondary antibody(SP-0023) and DAB(C-0010) staining



Blank control: 293T(blue). Primary Antibody:Rabbit Anti-VNN1 antibody(bs-10314R), Dilution: 5µg in 100 1µL 1X PBS containing 0.5% BSA; Isotype Control Antibody: Rabbit IgG(orange) ,used under the same conditions); Secondary Antibody: Goat anti-rabbit IgG-PE(white blue), Dilution: 1:200 in 1 X PBS containing 0.5% BSA. Protocol The cells were washed twice with phosphate-buffered saline (PBS). The cells were then incubated in 1 X PBS containing 0.5% BSA + 1 0% goat serum (15 min) to block non-specific protein-protein interactions followed by the antibody (bs-10314R, 5µg /1x10^6 cells) for 30 min on ice. The secondary antibody used was Goat Antirabbit IgG/PE antibody at 1/200 dilution for 30 min on ice. Acquisition of 20,000 events was performed.

Sheep,



Blank control: 293T. Primary Antibody (green line): Rabbit Anti-VNN1 antibody (bs-10314R) Dilution: 1µg /10^6 cells; Isotype Control Antibody (orange line): Rabbit IgG . Secondary Antibody : Goat anti-rabbit IgG-PE Dilution: 1µg /test. Protocol The cells were incubated in 5%BSA to block non-specific protein-protein interactions for 30 min at at room temperature .Cells stained with Primary Antibody for 30 min at room temperature. The secondary antibody used for 40 min at room temperature. Acquisition of 20,000 events was performed.