

Full-Length

BSND (Human) Recombinant Protein (P01)

Catalog # H00007809-P01

Size 50 ug

Specification

Product Description	Human BSND full-length ORF (1 a.a. - 320 a.a.) recombinant protein with GST-tag at N-terminal.
Sequence	MADEKTRIGFIVLGLFLLALGTFLMSHDRPQVYGTFYAMGSVMVIGGIWSMCQCYPKITFVPADS DFQGILSPKAMGLENGLAAEMKSPSPQPPYVRLWEEAAYDQSLPDFSHIQMKVMSYSEDHRSL LAPEMGQPKLGTSDGGEGGPGDVQAWMEAAVVIHKGSDESEGERRLTQSWPGPLACPQGPA PLASFQDDLDMSSEGSSPNASPHDREEACSPQQEPQGCRCPLDRFQDFALIDAPLTLEDEPQE GQQWEIALPNNWQRYPRTKVEEKEASDTGGEEPEKEEEDLYGLPDGAGDLLPDKELGFEPDT QG
Host	Wheat Germ (in vitro)
Theoretical MW (kDa)	61.6
Interspecies Antigen Sequence	Mouse (70); Rat (71)
Preparation Method	in vitro wheat germ expression system
Purification	Glutathione Sepharose 4 Fast Flow
Storage Buffer	50 mM Tris-HCl, 10 mM reduced Glutathione, pH=8.0 in the elution buffer.
Storage Instruction	Store at -80°C. Aliquot to avoid repeated freezing and thawing.
Note	Best use within three months from the date of receipt of this protein.

Applications

- Enzyme-linked Immunoabsorbent Assay
- Western Blot (Recombinant protein)
- Antibody Production

- Protein Array

Gene Info — BSND

Entrez GeneID	7809
GeneBank Accession#	AK129999.1
Gene Name	BSND
Gene Alias	BART, MGC119283, MGC119284, MGC119285
Gene Description	Bartter syndrome, infantile, with sensorineural deafness (Barttin)
Omim ID	602522 606412
Gene Ontology	Hyperlink
Gene Summary	This gene encodes an essential beta subunit for CLC chloride channels. These heteromeric channels localize to basolateral membranes of renal tubules and of potassium-secreting epithelia of the inner ear. Mutations in this gene have been associated with Bartter syndrome with sensorineural deafness. [provided by RefSeq]
Other Designations	OTTHUMP00000009804 barttin

Disease

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