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Lieferung & Zahlungsart

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Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

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Datasheet

CST3 (Human) Recombinant Protein (P01)

Catalog Number: H00001471-P01

Regulation Status: For research use only (RUO)

Product Description: Human CST3 full-length ORF (AAH13083, 1 a.a. - 146 a.a.) recombinant protein with GST-tag at N-terminal.

Sequence:

MAGPLRAPLLLLLAILAVALAVSPAAGSSPGKPPRLVGG
PMDASVEEEGVRRALDFAVGEYKASNDMYHSRALQ
VVRARKQIVAGVNYFLDVELGRTTCTKTQPNLDNCPF
HDQPHLKRKAFCSFQIYAVPWQGTMTLSKSTCQDA

Host: Wheat Germ (in vitro)

Theoretical MW (kDa): 41.69

Applications: AP, Array, ELISA, WB-Re

(See our web site product page for detailed applications information)

Protocols: See our web site at

<http://www.abnova.com/support/protocols.asp> or product page for detailed protocols

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.

Entrez GeneID: 1471

Gene Symbol: CST3

Gene Alias: ARMD11, MGC117328

Gene Summary: The cystatin superfamily encompasses proteins that contain multiple cystatin-like sequences. Some of the members are active cysteine protease inhibitors, while others have lost or perhaps never

acquired this inhibitory activity. There are three inhibitory families in the superfamily, including the type 1 cystatins (stefins), type 2 cystatins and the kininogens. The type 2 cystatin proteins are a class of cysteine proteinase inhibitors found in a variety of human fluids and secretions, where they appear to provide protective functions. The cystatin locus on chromosome 20 contains the majority of the type 2 cystatin genes and pseudogenes. This gene is located in the cystatin locus and encodes the most abundant extracellular inhibitor of cysteine proteases, which is found in high concentrations in biological fluids and is expressed in virtually all organs of the body. A mutation in this gene has been associated with amyloid angiopathy. Expression of this protein in vascular wall smooth muscle cells is severely reduced in both atherosclerotic and aneurysmal aortic lesions, establishing its role in vascular disease. [provided by RefSeq]