PRODUCT DATASHEET

Connective Tissue Growth Factor Human HEK293

Cat. No.: RD172035100-HEK
Type: Recombinant protein
Size: 0.1 mg
Source: HEK293
Species: Human

Description
Total 329 AA, UniProt P29279 (Gln27-Ala349). MW: 36 kDa (calculated). C-terminal 6×His tag. Protein identity confirmed by LC-MS/MS.

Other names
Hypertrophic chondrocyte-specific protein 24, Insulin-like growth factor-binding protein 8, IGF-binding protein 8, IGFBP-8, IBP-8, CCN family member 2, CTGF, CCN2, HCS24

Introduction to the molecule
Connective Tissue Growth Factor belongs to the CCN family of proteins. The CCN family presently consists of six members in human also known as: Cyr61 (Cystein rich 61), CTGF (Connective Tissue Growth Factor), Nov (Nephroblastoma Overexpressed gene), WISP-1, 2 and 3 (Wnt-1 Induced Secreted Proteins). The CCN genes encode secreted proteins associated with the Extracellular Matrix (ECM) and cell membrane. CCN proteins are matricellular proteins which are involved in the regulation of various cellular functions including: proliferation, differentiation, survival, adhesion and migration. They are expressed in derivatives of the three embryonic sheets and are implicated in the development of kidney, nervous system, muscle, bone marrow, cartilage and bone. During adulthood, they are implicated in wound healing, bone fracture repair, and pathologies such as: fibrosis, vascular ailments and tumourigenesis. Full length secreted CCN proteins can show an antiproliferative activity, whereas truncated isoforms are likely to stimulate proliferation and behave as oncogenes. The full-length protein consists of four modules: - Module I shares partial identity with the N-terminal part of the Insulin-like Growth Factor Binding Proteins (IGFBPs). - Module II includes a stretch of 70amino acid residues – which shares sequence identity with the Von Willebrand Factor Type C repeat (VWC). - Module III contains sequences sharing identity with the Thrombospondin type 1 repeat (TSP1) (WSXCSXXCG), which is thought to be implicated in the binding of sulfated glycoconjugates and to be important for cell adhesion. - Module IV, also designated CT, is encoded by exon 5. It is the least conserved one of the four domains at the level of nucleotide sequence, but it appears to be critical for several of the biological functions attributed to the CCN proteins. Module IV resembles the CT domain of several extracellular protein including, Von Willebrand’s fac-tor and mucins. Sequence similarities to heparin-binding motifs are also found within this domain. Proteolysis of the secreted full-length CCN proteins that has been reported in the case of CCN2 and CCN3 might result in the production of CCN-derived peptides with high affinity for ligands that full-length CNN proteins bind only poorly. Amino-truncated CCN2 isoforms were biologically active whereas no specific biological activity has been attributed to the truncated CCN3. Although the molecular processes underlying the production of these secreted isoforms is presently unknown, it is important to note that proteolysis occur at the same amino acid residues in both CCN2 and CCN3. An elevated expression of CCN2 has also been detected by Northern blotting in human invasive mammary ductal carcinomas, dermatofibromas, pyogenic granuloma, endothelial cells of angiolipomas and angioleiomomas, and in pancreatic tumours. A study performed with chondrosarcomas representative of various histological grades established that CCN2 expression was closely correlated with increasing levels of malignancy. In agreement with CCN2 playing a role in brain tumour angiogenesis, immunocytchemistry studies indicated that both glioblastoma tumour cells and proliferating endothelial cells stained positive for CCN2. In astrocytomas, CCN2 expression was particularly elevated in high-grade tumours, with a marked effect of CCN2 on cell proliferation. Downregulation of CCN2 expression in these cells was associated with a growth arrest at the G1/S transition while over-expression of CCN2 induced a two-fold increase of the number of cells in the G1 phase. Gene profiling analysis allowed to identify a set of about 50 genes whose expression might account for the proliferative activity of CCN2 in these cells. CCN2 was seen in a higher proportion of mononuclear cells of patients with acute lymphoblastic leukemia.

Research topic
Animal studies, Others
Amino Acid sequence
QNCSGPRCP DEAPRCPAG VSLVLGCGGC CRVCAKQLGE LCTERDPDCP HKGLFCFHS PANKIGVCT AKDGPCICFG GTVYRSGESF QSSKYQCTC LDGAVGCPL CSMVRRLPSD DCPFPVRVKL PGKCCCEWVC DEPKDQTVVG PALARERLED TFGPDPTMIR ANCLVQTTAEW SASKTCGMG ISTRVTNDNA SCRLEKQSRM CMVRPCEADL EENIKGGKKE KFELSGCSTM KTYRAKCFGV CTDGRCCCTPH RRTTRLPVEFK CPDGEMKKN MMFICTACCH YNCRDNDIF ESLYRKMYG DMAHHHHHH

Purity
Purity as determined by densitometric image analysis: >90%

Endotoxin
< 0.1 EU/μg

Formulation:
Sterile filtered (0.2 μm) and frozen from 0.10 – 0.30mg/mL solution in 0.1M citrate buffer pH 4.7 containing 20% (w/v) glycerol.

Reconstituion:
Defrost at ambient temperature

Shipping
On ice. Upon receipt, store the product at the temperature recommended below.

Storage, Stability/Shelf Life
Store protein at –80°C. Protein remains stable until the expiry date when stored at –80°C. Avoid repeated freezing/thawing cycles. The thawed protein can be stored at 4 °C for a week.

Quality control
BCA to determine quantity of the protein.

SDS PAGE to determine purity of the protein.

LAL to determine quantity of endotoxin.

Applications
Cell culture and/or animal studies, ELISA, Western blotting
12% SDS-PAGE separation of Human CTGF HEK293:
1. M.W. marker – 14, 21, 31, 45, 66, 97 kDa
2. reduced and heated sample, 2.5μg / lane
3. non-reduced and non-heated sample, 2.5μg / lane