# abcam

## Product datasheet

# Recombinant Human Ext2 protein ab158395

### 1 Image

Description		
Product name	Recombinant Human Ext2 protein	
Expression system	Wheat germ	
Protein length	Full length protein	
Animal free	No	
Nature	Recombinant	
Species	Human	
Sequence		MCASVKYNIRGPALIPRMKTKHRIYYTLFSIVLLGLIATGMFQ FWPHSI ESSNDWNVEKRSIRDVPVVRLPADSPIPERGDLSCRMHT CFDVYRCGFNP KNKIKVYTALKKYVDDFGVSVSNTISREYNELLMAISDSDY YTDDINRA CLFVPSIDVLNQNTLRIKETAQAMAQLSRWDRGTNHLLFN MLPGGPPDYN TALDVPRDRALLAGGGFSTWTYRQGYDVSIPVYSPLSAEV DLPEKGPGPR QYFLLSSQVGLHPEYREDLEALQVKHGESVLVLDKCTNLS EGVLSVRKRC HKHQVFDYPQVLQEATFCVVLRGARLGQAVLSDVLQAGC VPVVIADSYIL PFSEVLDWKRASVVVPEEKMSDVYSILQSIPQRQIEEMQR QARWFWEAYF QSIKAIALATLQIINDRIYPYAAISYEEWNDPPAVKWGSVSN PLFLPLIP PQSQGFTAIVLTYDRVESLFRVITEVSKVPSLSKLLVVWNN QNKNPPEDS LWPKIRVPLKVVRTAENKLSNRFFPYDEIETEAVLAIDDDII MLTSDELQ FGYEVWREFPDRLVGYPGRLHLWDHEMNKWKYESEWTN EVSMVLTGAAFY HKYFNYLYTYKMPGDIKNWVDAHMNCEDIAMNFLVANVTG KAVIKVTPRK

Amino acids	1 to 718
Tags	GST tag N-Terminus

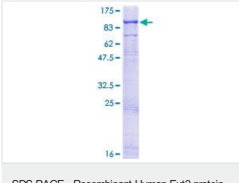
#### Specifications

Our <u>Abpromise guarantee</u> covers the use of ab158395 in the following tested applications.

The application notes include recommended starting dilutions; optimal dilutions/concentrations should be determined by the end user.

Applications	ELISA
	Western blot
Form	Liquid
Additional notes	

Preparation and Storage		
Stability and Storage	Shipped on dry ice. Upon delivery aliquot and store at -80°C. Avoid freeze / thaw cycles. pH: 8.00 Constituents: 0.31% Glutathione, 0.79% Tris HCI	
General Info		
Function	Glycosyltransferase required for the biosynthesis of heparan-sulfate. The EXT1/EXT2 complex possesses substantially higher glycosyltransferase activity than EXT1 or EXT2 alone. Appears to be a tumor suppressor.	
Tissue specificity	Ubiquitous.	
Pathway	Protein modification; protein glycosylation.	
Involvement in disease	<ul> <li>Defects in EXT2 are a cause of hereditary multiple exostoses type 2 (EXT2) [MIM:133701]. EXT is a genetically heterogeneous bone disorder caused by genes segregating on human chromosomes 8, 11, and 19 and designated EXT1, EXT2 and EXT3 respectively. EXT is a dominantly inherited skeletal disorder primarily affecting endochondral bone during growth. The disease is characterized by formation of numerous cartilage-capped, benign bone tumors (osteocartilaginous exostoses or osteochondromas) that are often accompanied by skeletal deformities and short stature. In a small percentage of cases exostoses have exhibited malignant transformation resulting in an osteosarcoma or chondrosarcoma. Osteochondromas development can also occur as a sporadic event.</li> <li>Defects in EXT2 are the cause of Potocki-Shaffer syndrome (PSS) [MIM:601224]. It is a contiguous gene syndrome due to proximal deletion of chromosome 11p11.2, including EXT2 and ALX4.</li> </ul>	
Sequence similarities	Belongs to the glycosyltransferase 47 family.	
Cellular localization	Endoplasmic reticulum membrane. Golgi apparatus membrane. The EXT1/EXT2 complex is localized in the Golgi apparatus.	



SDS-PAGE - Recombinant Human Ext2 protein (ab158395)

Please note: All products are "FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES"

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- Response to your inquiry within 24 hours
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- Extensive multi-media technical resources to help you
- We investigate all quality concerns to ensure our products perform to the highest standards

If the product does not perform as described on this datasheet, we will offer a refund or replacement. For full details of the Abpromise, please visit <u>https://www.abcam.com/abpromise</u> or contact our technical team.

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