

Recombinant Viral EBOV GP (Mucin Domain Deleted)

Catalog Number: 10585-EB

DESCRIPTION				
Source	Human embryonic kidney cell, HEK293-derived viral EBOV GP protein			
	Ebola Virus GP1 (Ile33-Val311) (Thr42Val, Thr230Val) Accession # NP_0066246.1	Ebola Virus GP2 (Thr464-Asp632) Accession # NP_066246.1	ннннн	
	N-terminus		C-terminus	
N-terminal Sequence Analysis	Ile33 & Glu502			
Predicted Molecular Mass	51 kDa			
SPECIFICATIONS				
SDS-PAGE	63-75 kDa under non-reducing conditions			

SDS-PAGE	63-75 kDa, under non-reducing conditions Measured by its binding ability in a functional ELISA. When Recombinant Viral EBOV GP (mucin domain deleted, Catalog # 10585-EB) is immobilized at 2 μg/mL (100 μL/well), the concentration of Recombinant Human CLEC10A/CD301 (Catalog # 4888-CL) binds with an ED ₅₀ of 1-10 ng/mL.	
Activity		
Endotoxin Level	<0.10 EU per 1 µg of the protein by the LAL method.	
Purity	>95%, by SDS-PAGE visualized with Silver Staining and quantitative densitometry by Coomassie® Blue Staining.	
Formulation	Lyophilized from a 0.2 µm filtered solution in PBS. See Certificate of Analysis for details.	

PREPARATION AND STORAGE			
Reconstitution	Reconstitute at 500 μg/mL in PBS.		
Shipping	The product is shipped at ambient temperature. Upon receipt, store it immediately at the temperature recommended below.		
Stability & Storage	Use a manual defrost freezer and avoid repeated freeze-thaw cycles.		
	 12 months from date of receipt, -20 to -70 °C as supplied. 		
	1 month, 2 to 8 °C under sterile conditions after reconstitution.		

• 3 months, -20 to -70 °C under sterile conditions after reconstitution.



When Recombinant Viral EBOV GP (Mucin Domain Deleted) Protein (Catalog # 10585-EB) is immobilized at 2 μ g/mL (100 μ L/well), the concentration of Recombinant Human CLEC10A/CD301 Protein (Catalog # 4888-CL) binds with an ED₅₀ of 1-10 ng/mL.

Rev. 12/2/2020 Page 1 of 2



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BACKGROUND

The GP glycoprotein encoded by the genome of Ebola family viruses is a critical molecule for the pathogenicity of Ebolavirus hemorrhagic viruses (1, 2). It is processed into distinct forms for virus capsule or cell surface presentation or release from virus infected cells. The GP precursor protein is cleaved by furin at a multibasic site to yield a 140 kDa N-terminal fragment (GP1) and a 26 kDa C-terminal fragment (GP2) which remain disulfide linked (3). GP1 is entirely extracellular while GP2 is a transmembrane protein (4). Heterodimers of GP1-GP2 can further associate into trimers (5). GP expressed on virus infected cells are beshed by TACE mediated cleavage, liberating a disulfide linked complex of soluble GP1 and truncated GP2 (4-6). GP binds to multiple C-type lectins on target cell surfaces, including CLEC10A/MGL, DC-SIGNR (7-9). Following internalization, GP1 is cleaved by Cathepsin B and Cathepsin L and then interacts with Niemann-Pick C1 (NPC1) in the endosomal membrane (10-12).

References:

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Rev. 12/2/2020 Page 2 of 2



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