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## 12-8207: Anti-Ebolavirus, GP (Clone EBOV-548)-Purified No Carrier Protein

Clonality: Monoclonal Clone Name: EBOV-548 Isotype: Human IgG1

#### Description

Specificity: EBOV-548 activity is directed against the glycan cap of the GP1 subunit of Zaire ebolavirus (EBOV) and Bundibugyo ebolavirus (BDBV) glycoprotein.

Antigen Distribution: Ebola virus glycoprotein is a surface protein expressed on the virus envelope.

Background: Ebola virus is a member of the Filoviridae family that causes severe disease in humans with a mortality rate of 25-90%1. Three Ebola species are responsible for lethal outbreaks: Zaire ebolavirus (EBOV), Bundibugyo ebolavirus (BDBV), and Sudan ebolavirus (SUDV). The Ebola virus envelope contains a single surface glycoprotein (GP) which is responsible for viral attachment to the host cell, endosomal entry, and membrane fusion1. GP is composed of two subunits, GP1 and GP2. GP1 has a heavily glycosylated mucin-like domain and a glycan cap. GP2 contains the internal fusion loop, transmembrane domain, and stalk. GP is the major target of neutralizing monoclonal antibody (mAb) and vaccine design against Ebola virus1,2,3,4. EBOV-548 is a pan-EBOV-neutralizing mAb isolated from a survivor of the EBOV 2013-2016 outbreak2. Hybridomas were generated from human peripheral blood mononuclear cells. EBOV GP-reactive memory B cells were labeled with recombinant EBOV GP protein, purified by FACS, bulk expanded on NIH 3T3 cells, bulk fused with MFP-2 myeloma cells, and screened for neutralizing activity against live and/or recombinant EBOV, BDBV, and SUDV GP. EBOV-548 reacts to all three species GPs and neutralizes EBOV and BDBV. EBOV-548 recognizes intact but not cleaved GP. EBOV-548 was identified in a study designed to develop a cooperative two-antibody cocktail with EBOV-5202,4. When paired, EBOV-548 and EBOV-520 have synergistic activity for GP binding and virus neutralization2. This is achieved mechanically. EBOV-548 binds to the glycan cap in a manner that destabilizes the GP trimer, displacing the ?17-?18 loop. As a result, the glycan cap is pulled back, facilitating EBOV-520 binding and enhancing neutralization. The cooperative effect is dependent on EBOV-548 concentration. When administered together in vivo, EBOV-548 potentiates protection by EBOV-520 against SUDV, with 50% of mice surviving treatment, while only 10% survive with EBOV-520 alone.

### **Product Info**

| Amount :<br>Purification : | 1 mg / 250μg<br>Purity: >=90% monomer by analytical SEC and SDS-Page<br>Preparation: Recombinant antibodies are manufactured in an animal free facility using only in<br>vitro protein free cell culture techniques and are purified by a multi-step process including the<br>use of protein A or G to assure extremely low levels of endotoxins, leachable protein A or<br>aggregates.   |
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| Content :                  | Concentration: >=1.0 mg/ml<br>Formulation: This recombinant monoclonal antibody is aseptically packaged and formulated in<br>0.01 M phosphate buffered saline (150 mM NaCl) PBS pH 7.2 - 7.4 with no carrier protein,<br>potassium, calcium or preservatives added. Due to inherent biochemical properties of<br>antibodies, certain products may be prone to precipitation over time. Precipitation may be<br>removed by aseptic centrifugation and/or filtration. |
| Storage condition :        | This antibody may be stored sterile as received at 2-8°C for up to one month. For longer term storage, aseptically aliquot in working volumes without diluting and store at $<=$ -70°C.?Avoid Repeated Freeze Thaw Cycles.  |

# **Application Note**

#### B, ELISA, EM, FA, N