

PRODUCT INFORMATION

Clone ID DMC480 **Target** DDR1

CAK; EDDR1; NEP; NTRK4; PTK3A; RTK6; TRKE; **Synonyms**

MCK-10; HGK2; CD167a

Host Species

Anti-DDR1 antibody(DMC480); IgG1 Chimeric **Description**

mAb Delivery In Stock **Uniprot ID** 008345

IgG type Rabbit/Human Fc chimeric IgG1

Clonality Monoclonal Reactivity Human **Applications** Flow Cyt

Recommended

Flow Cyt 1:100 **Dilutions**

Purified from cell culture supernatant by affinity **Purification**

chromatography

Lyophilized from sterile PBS, pH 7.4. Normally 5 % - 8% trehalose is added as protectants before lyophilization. Please see Certificate of Analysis Formulation & Reconstitution

for specific instructions of reconstitution. Store at -20°C to -80°C for 12 months in lyophilized form. After reconstitution, if not

intended for use within a month, aliquot and store Storage & Shipping at -80°C (Avoid repeated freezing and thawing). Lyophilized proteins are shipped at ambient

temperature.

Receptor tyrosine kinases play a key role in the communication of cells with their

microenvironment. These kinases are involved in the regulation of cell growth; differentiation and metabolism. The protein encoded by this gene belongs to a subfamily of tyrosine kinase receptors with homology to Dictyostelium discoideum protein discoidin I in their

extracellular domain; and that are activated by various types of collagen. Expression of this Background protein is restricted to epithelial cells; particularly

in the kidney; lung; gastrointestinal tract; and brain. In addition; it has been shown to be significantly overexpressed in several human tumors. Alternatively spliced transcript variants encoding different isoforms have been described for this gene. [provided by RefSeq; Feb 2011]

Usage Research use only Conjugate Unconjugated

All DIMA recombinant antibodies are genuinely generated by DIMA Biotech. They are all under patent application. Any protein sequencing or reverse engineering attempt is prohibited. We are

actively scrutinizing all patent application to

ensure no IP infringement.

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DIMA Disclaimer

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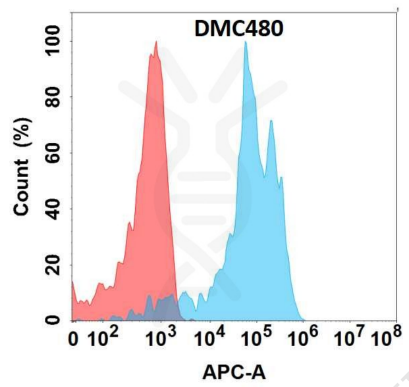


Figure 1. Flow cytometry analysis with Anti-DDR1 (DMC480) on HEK293 cells transfected with human DDR1 (Blue histogram) or HEK293 transfected with irrelevant protein (Red histogram).

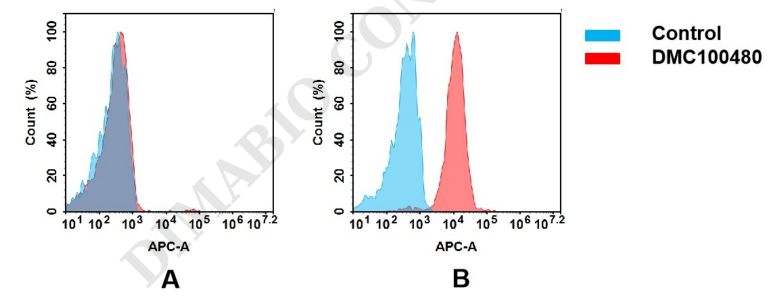


Figure 2. Flow cytometry analysis of antigen binding of anti-human DDR1 mAb(DMC100480).
(A) DMC100480 does not bind to jurkat cells that do not express DDR1.
(B) A clear peak shift of DMC100480 was seen compared to the control when incubated with DDR1-expressing SNU-5 cells, indicating strong binding of DMC100480 to DDR1. Antibodies were incubated at 5 μg/mL.









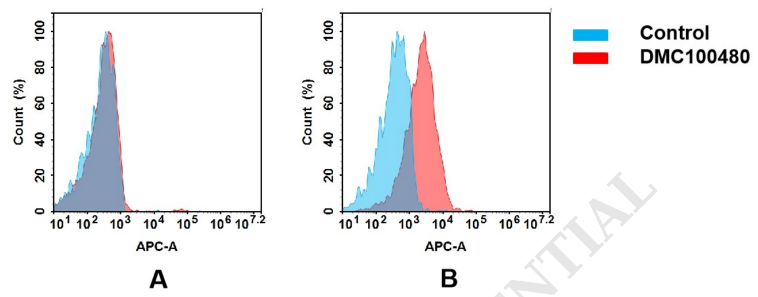


Figure 3. Flow cytometry analysis of antigen binding of anti-human DDR1 mAb(DMC100480). (A) DMC100480 does not bind to jurkat cells that do not express DDR1. (B) A clear peak shift of DMC100480 was seen compared to the control when incubated with DDR1-expressing MCF-7 cells, indicating strong binding of DMC100480 to DDR1. Antibodies were incubated at 5 μ g/mL.



