# anti-ZKSCAN2-RAB-C205



# **Applications**

Competition ELISA	Western Blot	SPR	Spiked IP	Immunofluorescence	IP-MS	ChIP
Pass			Pass	Pass		

<sup>\*</sup>rAb has been tested for the following applications. See below for the experimental details.

## **Antibody information**

rAb ID: anti-ZKSCAN2-RAB-C205

**Description:** recombinant Fab fragment obtained by recombinant antibody (rAb) phage display recognizing *ZKSCAN2* protein under non-denaturing conditions; specificity and affinity tested.

**Binder type:** rAb **Isotype:** IgG1 **Species:** *Homo sapiens* **Produced in:** *E. coli* **rAb tags:** Avi-tag; no tag **Specificity:** reacts with *Homo sapiens* ZKSCAN2 **Epitope:** binds to folded domain amino acids 36-131

Storage conditions: short term - store at n 4°C (over 6 months), long term - PBS -20°C or -80°C

Link: http://recombinant-antibodies.org/binders/anti-ZKSCAN2-RAB-C205

## **Antigen information**

**Protein Name:** Zinc finger protein with KRAB and SCAN domains 2 **HGNC Symbol:** ZKSCAN2 **HGNC ID:** 25677 **Species:** *Homo sapiens* 

UniProt AC: Q63HK3 UniProt KB: ZKSC2 HUMAN

**Protein Sequence:** 

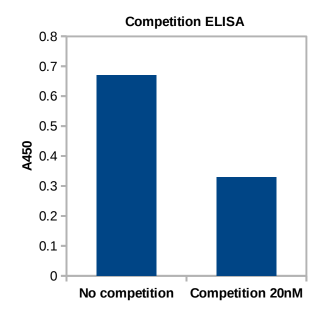
MSGLNDIFEAQKIEWHEHHHHHHENLYFQSHMEGSDSSETFRKCFRQFCYEDVTGPHEAFSKLWELCCRWLKPEMRSKE QILELLVIEQFLTILPEKIQAWAQKQCPQSGEEAVALVVHLEKETGRLR

**Vector Type:** pET15Avi6HT\_NESG **Vector Link:** http://beta.labgeni.us/registries/DNASU/pET15Avi6HT\_NESG/

Protein Sequence Position: 36-131 Antigen source: E. coli Source Lab: Rutgers Source Lab ID: HR8296A.002

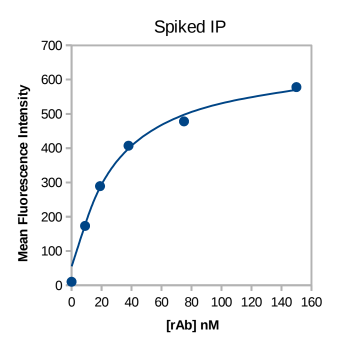
**Description:** affinity purified recombinant protein

## Validation data



**Single point competition phage ELISA** Plot represents specific binding of a target to the rAb-phage in solution (right bar) in comparison to binding to the target immobilized on the plate surface (left bar). Experimental conditions were calibrated to capture binders with dissociation Constant ( $K_D$ ): 20nM or lower.

Experimental Conditions: Culture supernatants containing rAb-phage were diluted five-fold in phosphate-buffered saline, 0.5% (w/v) BSA, 0.1% (v/v) Tween 20 either with or without soluble antigen competitor at 20 nM. After 1 h incubation at room temperature, the mixtures were transferred to neutravidin coated plates preloaded with 50  $\mu L$  of 20 nM biotinylated antigen and incubated for 15 min. The plates were washed with phosphate-buffered saline, 0.05% (v/v) Tween 20 and incubated for 30 min with horse radish peroxidase/anti-M13 antibody conjugate (1:5000 dilution). The plates were washed, developed with 3,3',5,5'-Tetramethyl-benzidine/H<sub>2</sub>O<sub>2</sub> peroxidase substrate (Thermo Scientific), quenched with 1M H<sub>3</sub>PO<sub>4</sub>, and the absorbance at 450 nm (A450) was determined.



**Spiked IP** Tritration curve of rAb against antigen of interest. The  $K_D$  values were obtained by the least-squares fitting of fluorescence saturation data.

Dissociation Constant (K<sub>D</sub>): 26.4 ± 3.3 nM

#### **Experimental Conditions:**

**Spiked IP:** Antigen was immobilized to M280 Dynabeads. A rAb, 50 nM, was pulled down from high salt AFC buffer with or without HEK293 lysate (OD280 ~10). Beads were washed with low salt AFC buffer, and the captured antibody was quantified with an anti-Fab fluorophore labeled antibody on a flow cytometer.

**Affinity Measurement:** Antigen was immobilized to M280 Dynabeads and incubated with a rAb, in varying concentration (100 nM down to 1 nM in three-fold dilutions). Beads were washed with BSET/BSA and quantified.

#### **Buffers:**

High salt AFC buffer: 10 mM Tris-HCl, pH 7.9, 420 mM NaCl, 0.1% NP-40

Low salt AFC buffer: 10 mM Tris-HCl, pH 7.9, 100 mM NaCl, 0.1% NP-40

PBSE/BSA: 20 mM Na2HPO4, pH 7.5, 150 mM NaCl, 1 mM EDTA. 0.5% BSA

PBSET/BSA: PBSE/BSA + 0.1% Tween-20

Immunofluorescence:
Status: Pass
Experimental Conditions: <a href="http://recombinant-antibodies.org/protocols/immunofluorescence">http://recombinant-antibodies.org/protocols/immunofluorescence</a>
IP-MS – immunoprecipitation for mass spectrometric analysis:
Status:
Experimental Conditions: http://recombinant-antibodies.org/protocols/IP-MS
ChIP – chromatin immunoprecipitation:
Status:
Experimental Conditions: Pending
IP – immunoprecipitation:
Status:
Experimental Conditions: Pending
SP Elisa:
Status:
Experimental Conditions: <a href="http://recombinant-antibodies.org/protocols/ELISA-IC50-EC50-direct-coating">http://recombinant-antibodies.org/protocols/ELISA-IC50-EC50-direct-coating</a>

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