#### For Research Use Only

# Phospho-CHEK2 (Thr68) Polyclonal antibody



**Purification Method:** 

Catalog Number: 29012-1-AP

5 Publications

**Basic Information** 

Catalog Number: GenBank Accession Number: BC004207

Antigen affinity purification 29012-1-AP GeneID (NCBI): Recommended Dilutions: WB 1:500-1:2000

100ul, Concentration: 450 µg/ml by 11200

Source: CHK2 checkpoint homolog (S. pombe)

Rabbit Calculated MW: Isotype: 61 kDa IgG Observed MW:

65 kDa

**Applications** 

**Tested Applications:** 

WB, ELISA

**Cited Applications:** 

WB

Species Specificity:

Human **Cited Species:** human

Positive Controls:

WB: MMS treated PC-3 cells,

## **Background Information**

Serine/threonine-protein kinase Chk2 (CHEK2) is a serine/threonine kinase which is activated upon DNA damage and is implicated in pathways that govern DNA repair, cell cycle arrest or apoptosis in response to the initial  $damage.\,ATM\,phosphory lates\,CHEK2\,on\,T68.\,Phosphory lation\,on\,T68\,and\,subsequent\,full\,activation\,of\,CHEK2\,was$ shown to require priming phosphorylation on adjacent residues by Polo-like kinase 3 (PLK3) and the dualspecificity tyrosine and serine/threoninekinase TTK/hMPS1. Additionally TTK appears to phosphorylate T68. Phosphorylation of T68 promotes the binding of the N-terminal SQ/TQ-rich cluster of one CHEK2 molecule with the FHA domain of another CHEK2 molecule. (PMID: 28553140, PMID: 18004398, PMID: 33322746)

#### **Notable Publications**

Author	Pubmed ID	Journal	Application
Xin Wen	36249018	Front Oncol	WB
Zhili Xia	36185307	Front Oncol	WB
Chao Mei	35187743	Cell Prolif	WB

Storage

Storage:

Store at -20°C. Storage Buffer:

PBS with 0.02% sodium azide and 50% glycerol pH 7.3.

Aliquoting is unnecessary for -20°C storage

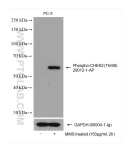
\*\*\* 20ul sizes contain 0.1% BSA

in USA), or 1(312) 455-8498 (outside USA)

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## Selected Validation Data



Non-treated PC-3 and MMS treated PC-3 cells were subjected to SDS PAGE followed by western blot with 29012-1-AP (Phospho-CHEK2 (Thr68) antibody) at dilution of 1:1000 incubated at room temperature for 4°C overnight. The membrane was stripped and re-blotted with GAPDH antibody as loading control.