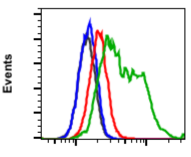
**Catalog:** #2116

Store at: -20ºC

For Research Use Only. Not For Use In Diagnostic Procedures.

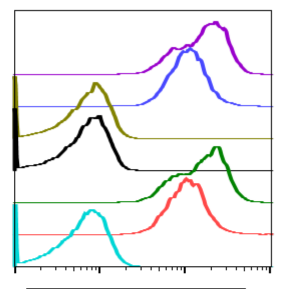
<b>Applications</b> Flow Cytometry,WB	<b>Detection</b> Anti-Rabbit IgG	<b>Clonality</b> Monoclonal	<b>lsotype</b> Rabbit lgGk
Format:	Unconjugated		
Cross Reactivity:	Predicted to work with mouse, rat, and other homologues		
Formulation:	1X PBS, 0.02% NaN3, 50% Glycerol, 0.1% BSA		
Preparation:	Protein A+G		
Reactivity:	Human,Mouse		
Recommended Usage:	$1\mu$ g/mL – 0.001 $\mu$ g/mL. It is recommended that the reagent be titrated for optimal performance for each application. See product image legends for additional information.		
Immunogen:	A synthetic phospho-peptide corresponding to residues surrounding Thr68 of human phospho Chk2		
Description:	Checkpoint kinase 2 (Chk2) plays a n DNA damage. Chk2 is initially inactiv Phosphorylation at Thr68 induces ho autophosphorylation within the kinas events within the auto-inhibitory loop phosphorylations, active dimers and substrates such as Cdc25C and BRCA missense variants have been found to breast and colon cancer. Constitutive found in many human cancer cell line	re in its monomeric, unpho modimerization, initiating se loop at Ser516 and photo o at Thr383 and Thr387. A monomers can then phos A1. In humans, Chk2 gener to be associated with incre ely phosphorylated Chk2 a	osphorylated form. rphorylation fter these phorylate tic deletion and eased risk of at Thr68 has been
References:	Ahn J, Urist M, and Prives C. (2004) DNA Repair. 3: 1039-1047.		



Phospho-CHK2 (Thr68)

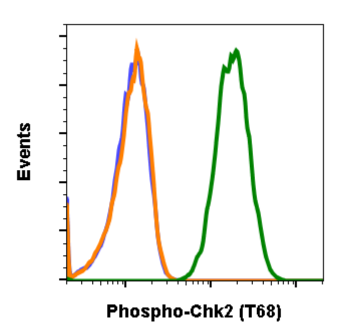
Flow cytometric analysis of Hela cells treated with lambda phosphatase (red) or treated with UV+TPA (green) using Phospho-CHK2 (Thr68) (D12) Rabbit mAb, at 0.01 ug/mL, CHK2T68-D12 #2116, or concentration-matched Rabbit (G9) mAb IgG Isotype Control #2141 for cells treated with lambda phosphatase (black) or treated with UV+TPA (blue).





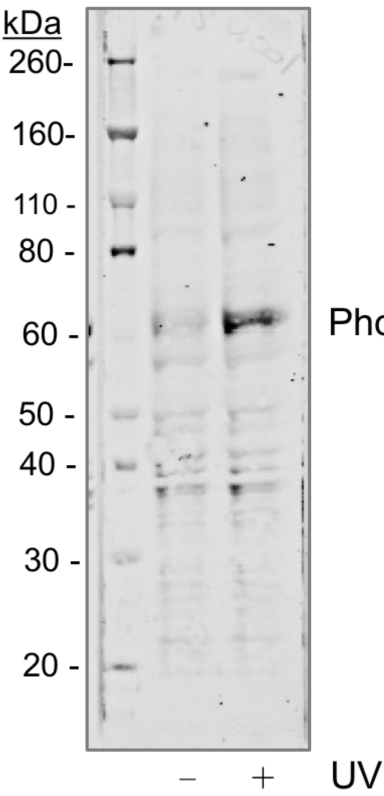
SampleID	Median : BL1-A
UVP D12 N	16792
Ctrl D12 N	10578
UVP D12 P	729
Ctrl D12 P	711
UVP D12	16290
Ctrl D12	10297
Ctrl 2' only	627

Peptide blocking flow cytometric analysis of HEK293T cells secondary antibody only negative control (light blue) or untreated (red) or UV/TPA-treated (green) or untreated and blocked with phospho-peptide (black) or UV/TPA and blocked with phospho peptide (gold) or untreated and blocked with non-phospho peptide (dark blue) or UV/TPA and blocked with non-phospho peptide (purple) using Phospho-Chk2 (T68) antibody Chk2T68-D12 0.1µg/mL. Cat. #2116.



Chk2T68-D12 recognizes basal phosphorylation levels in mouse cells. Flow cytometric analysis of 3T3 cells secondary antibody only (blue) or  $0.1 \mu g/mL$  of isotype control Cat. #2141 (orange) or of Phospho-Chk2 (T68) antibody Chk2T68-D12 (green) Cat. #2116.

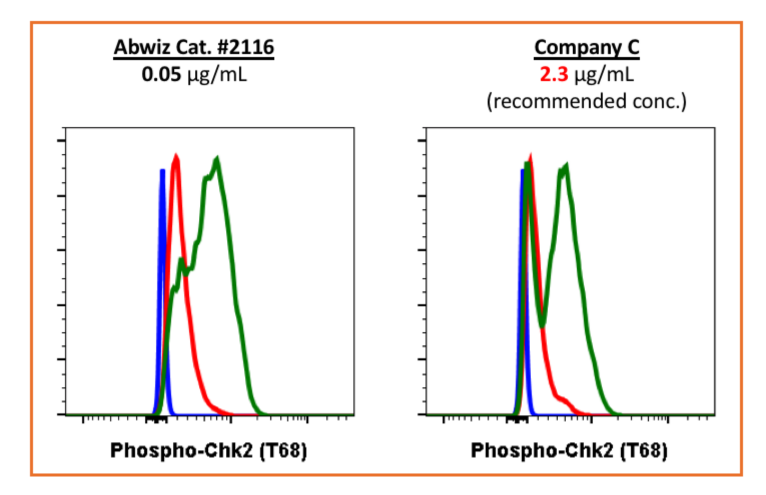




## Phospho-Chk2 (T68)

Western blot analysis of HEK293 cell extract untreated or treated with UV using Phospho Chk2(T68) antibody Chk2T68-D12. Cat. #2116.





Flow cytometric analysis of C6 cells secondary antibody only negative control (blue) or treated with imatinib (red) or with pervanadate (green) using 0.05  $\mu$ g/mL Phospho-Chk2 (T68) antibody Chk2T68-D12 (Abwiz Cat. #2116) or Company C antibody at 2.3  $\mu$ g/mL (manufacturer's recommended concentration).

