

## **BIK Ab**

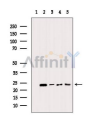
Cat.#: AF6428 Size: 100ul,200ul	Concn.: 1mg/ml Source: Rabbit	Mol.Wt.: 23kDa Clonality: Polyclonal
Application:	WB 1:500-1:2000 IHC 1:50-1:200 IF/ICC 1:100-1:500	
Reactivity:	Human, Mouse	
Purification:	The antiserum was purified by peptide affinity chromatography using SulfoLink™ Coupling Resin (Thermo Fisher Scientific).	
Specificity:	BIK Ab detects endogenous levels of total BIK.	
Immunogen:	A synthesized peptide derived from human BIK.	
Uniprot:	Q13323	
Description:	The protein encoded by this ger cellular and viral survival-promo and the Epstein-Barr virus in or cell death. Because its activity i of survival-promoting proteins, likely target for antiapoptotic pr	oting proteins, such as BCL2 der to enhance programed s suppressed in the presence this protein is suggested as a
Subcellular Location:	Endomembrane system. Around in cytoplasmic membranes.	the nuclear envelope, and
Similarity:	Intact BH3 motif is required by BIK, BID, BAK, BAD and BAX for their pro-apoptotic activity and for their interaction with anti-apoptotic members of the Bcl-2 family.	
Storage Condition and Buffer:	Rabbit IgG in phosphate buffered saline , pH 7.4, 150mM NaCl, 0.02% sodium azide and 50% glycerol.Store at -20 °C.Stable for 12 months from date of receipt.	



Western blot analysis of extracts from Hela, using BIK Ab. Lane 1 was treated with the blocking peptide.



Affinity Biosciences website:www.affbiotech.com order:order@affbiotech.com

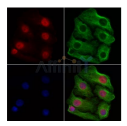


Western blot analysis of extracts from various samples, using BIK Ab.

Lane 1: Hybridoma cells treated with blocking peptide; Lane 2: Hybridoma cells; Lane 3: HeLa; Lane 4: Vero; Lane 5: HuvEc.



AF6428 at 1/100 staining Mouse muscle tissue by IHC-P. The sample was formaldehyde fixed and a heat mediated antigen retrieval step in citrate buffer was performed. The sample was then blocked and incubated with the Ab for 1.5 hours at 22°C. An HRP conjugated goat anti-rabbit Ab was used as the secondary.



AF6428 staining Hela cells by IF/ICC. The samples were fixed with PFA and permeabilized in 0.1% Triton X-100,then blocked in 10% serum for 45 minutes at 25°C. Samples were then incubated with primary Ab(AF6428 1:200) and mouse antibeta tubulin Ab(T0023 1:200) for 1 hour at 37°C. An AlexaFluor594 conjugated goat anti-rabbit IgG(H+L) Ab(Red) and an AlexaFluor488 conjugated goat anti-mouse IgG(H+L) Ab(Green) were used as the secondary Ab.



AF6428 staining HepG2 by IF/ICC. The sample were fixed with PFA and permeabilized in 0.1% Triton X-100,then blocked in 10% serum for 45 minutes at 25°C. The primary Ab was diluted at 1/200 and incubated with the sample for 1 hour at 37°C. An Alexa Fluor 594 conjugated goat anti-rabbit IgG (H+L) Ab, diluted at 1/600, was used as the secondary Ab.



AF6428 staining Jurkat cells by IF/ICC. The sample were fixed with PFA and permeabilized in 0.1% Triton X-100,then blocked in 10% serum for 45 minutes at 25°C. The primary Ab was diluted at 1/200 and incubated with the sample for 1 hour at 37°C. An Alexa Fluor 594 conjugated goat anti-rabbit IgG (H+L) Ab(Red), diluted at 1/600, was used as secondary Ab.

<code>IMPORTANT:</code> For western blot, incubate membrane with diluted primary Ab in 5% w/v milk , 1X TBS, 0.1% Tween®20 at 4°C with gentle shaking, overnight.

For Research Use Only. Not for use in diagnostic and therapeutic procedures. Not for resale without express authorization.