## DDB1 antibody

Catalog No: #38485

SAB Signalway Antibody

Orders: order@signalwayantibody.com Support: tech@signalwayantibody.com

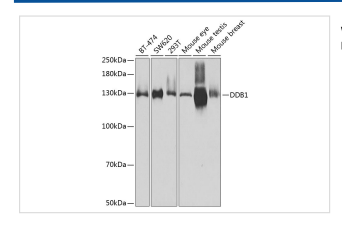
Package Size: #38485-1 50ul #38485-2 100ul

Description	
Product Name	DDB1 antibody
Host Species	Rabbit
Clonality	Polyclonal
Purification	Antibodies were purified by affinity purification using immunogen.
Applications	WB,IHC,IF
Species Reactivity	Human,Mouse,Rat
Specificity	The antibody detects endogenous level of total DDB1 protein.
Immunogen Type	Peptide
Immunogen Description	A synthetic peptide of human DDB1.
Target Name	DDB1
Other Names	XPE;DDBA;XAP1;XPCE;XPE-BF;UV-DDB1;
Accession No.	Swiss-Prot#: Q16531NCBI Gene ID: 1642
Uniprot	Q16531
GeneID	1642;
SDS-PAGE MW	127kd
Concentration	1.0mg/ml
Formulation	Supplied at 1.0mg/mL in phosphate buffered saline (without Mg2+ and Ca2+), pH 7.4, 150mM NaCl, 0.02%
	sodium azide and 50% glycerol.
Storage	Store at -20°C

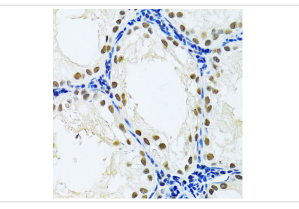
## Application Details

WB 1:500 - 1:1000IHC 1:50 - 1:100IF 1:50 - 1:100

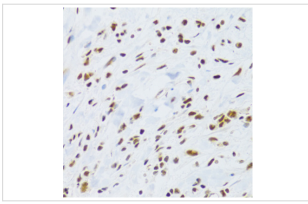
## **Images**



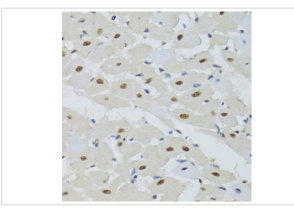
Western blot analysis of extracts of various cell lines, using DDB1 at 1:500 dilution.



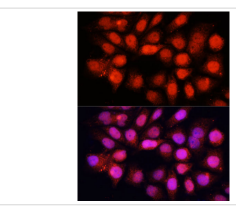
Immunohistochemistry of paraffin-embedded rat testis using DDB1 at dilution of 1:100 (40x lens).



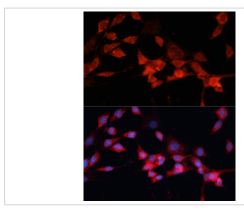
Immunohistochemistry of paraffin-embedded human gastric cancer using DDB1 at dilution of 1:100 (40x lens).



Immunohistochemistry of paraffin-embedded mouse heart using DDB1 at dilution of 1:100 (40x lens).



Immunofluorescence analysis of HeLa cells using DDB1 Polyclonal at dilution of 1:100 (40x lens). Blue: DAPI for nuclear staining.



Immunofluorescence analysis of NIH-3T3 cells using DDB1 Polyclonal at dilution of 1:100 (40x lens). Blue: DAPI for nuclear staining.

## Background

The protein encoded by this gene is the large subunit (p127) of the heterodimeric DNA damage-binding (DDB) complex while another protein (p48) forms the small subunit. This protein complex functions in nucleotide-excision repair and binds to DNA following UV damage. Defective activity of this complex causes the repair defect in patients with xeroderma pigmentosum complementation group E (XPE) - an autosomal recessive disorder characterized by photosensitivity and early onset of carcinomas. However, it remains for mutation analysis to demonstrate whether the defect in XPE patients is in this gene or the gene encoding the small subunit. In addition, Best vitelliform mascular dystrophy is mapped to the same region as this gene on 11q, but no sequence alternations of this gene are demonstrated in Best disease patients. The protein encoded by this gene also functions as an adaptor molecule for the cullin 4 (CUL4) ubiquitin E3 ligase complex by facilitating the binding of substrates to this complex and the ubiquitination of proteins.

Note: This product is for in vitro research use only