

## Anti-VRK1 (human) antibody, mouse monoclonal (5D1)

Product code 7	71-600
	100 μg
	20%
	1.0 mg/ml
<b>Buffer</b> P	PBS- with 50% glycerol
Purity P	Purified IgG fraction with protein A from hybridoma cell culture medium.
Immunogen S	Synthetic peptide corresponding to N-terminus of human VRK1,
N	MPRVKAAQAGRQSSAKRHL-C
Isotype	Mouse IgG1κ
Reactivity H	Human VRK1 protein. Not tested with other species.
Special notes N	N/A
Application 1	. Western blotting (1/200~1/1,000 dilution). Use of highly sensitive
	chemiluminescence reagents such as Lumi-Light Plus (Roche) or
	ImmunoStar®LD (Wako, Tokyo) are recommended.
2	2. Immunoprecipitation (assay dependent)
3	3. Immunofluorescence staining (1/100 dilution)
4	1. Immunohistochemistry (assay dependent)
5	5. ELISA (assay dependent)
Background T	The VRK1 gene encodes serine/threonine kinase VRK1 (Vaccinia-Related Kinase 1;
3	396 aa, 45.5 kDa) which is involved in Golgi disassembly during the cell cycle
fc	following phosphorylation by PLK3 during mitosis, and required to induce Golgi
fr	ragmentation. It acts by mediating phosphorylation of a downstream target protein
Γ'	Thr-18' of p53/TP53 and may thereby prevent the interaction between p53/TP53
a	and MDM2. It also phosphorylates casein and histone H3. Phosphorylation of the
В	BANF1 gene product disrupts its ability to bind DNA, reduces its binding to LEM
d	lomain-containing proteins and causes its relocalization from the nucleus to the
c.	ytoplasm.
Iı	nvolvement in disease Defects in VRK1 are the cause of pontocerebellar
h	hypoplasia type 1A (PCH1A); also called pontocerebellar hypoplasia with infantile
sj	spinal muscular atrophy or pontocerebellar hypoplasia with anterior horn cell
d	lisease. PCH1A is characterized by an abnormally small cerebellum and brainstem,
Ce	entral and peripheral motor dysfunction from birth, gliosis and anterior horn cell
d	legeneration resembling infantile spinal muscular atrophy.
Data Link S	7 : P + 000000 H
	SwissProt: Q99986 Human , Entrez Gene: 7443 Human
Please note: All product	ts are FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC



Data Images: 71-600 Anti-VRK1 (human) antibody, mouse monoclonal (5D1)

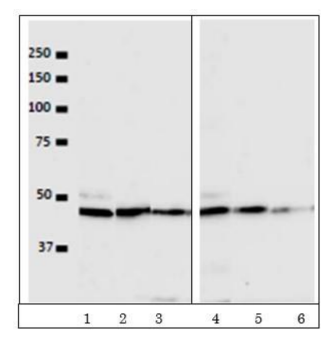


Fig.1 Western blot detection of VRK1 in the crude extracts of human cells. Lanes 1, 2, 3; HeLa cell extract (5x10<sup>4</sup> cells) with antibody dilutions at 1/100, 1/500, 1/1000. Lanes 4, 5, 6; U2OS cell extract (5x10<sup>4</sup> cells) with the antibody dilutions at 1/100, 1/500, 1/1,000. As secondary antibody, Alexa488 goat anti-mouse IgG was used. ImmunoStar®LD (Wako, Tokyo) was used as chemiluminescence reagent and images were taken with BIO-RAD ChemiDocXRS.

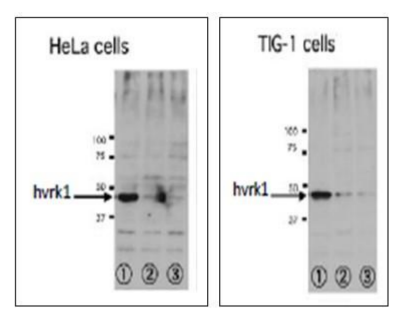


Fig.2. Inhibition of VRK1 expression in human cells treated by RNAi. specific to VRK1.

Lane 1; Luciferase RNAi (control). Lane2; VRK1-1 RNAi. Lane 3; VRK1-2 RNAi.. Antibody at 1/500 dilution. Lumi-Light Plus (Roche) was used as chemiluminescence reagent. Extracts from 5x10<sup>4</sup> cells.



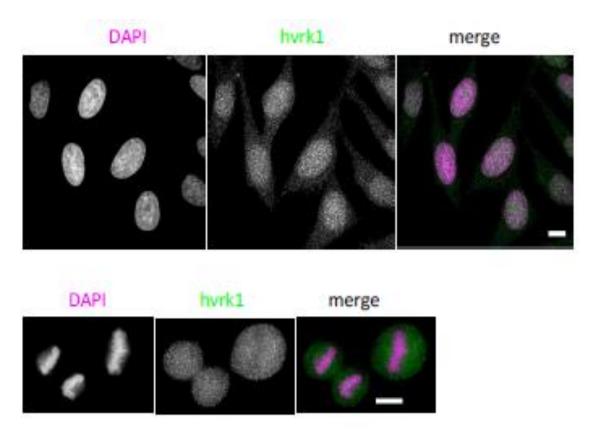


Fig.3. Immunoflorescence staining of VRK1 in HeLa cells, PA fixed.

Top; Interphase cells were fixed with paraformaldehyde and stained with the anti-human VRK1 antibody (hvrk1) at 1/100 dilution (center), DNA was stained with DAPI (left) and two images were merged (right; merge).

Bottom; Metaphase cells. At metaphase, VRK1 dots were solely detected in nuclei.



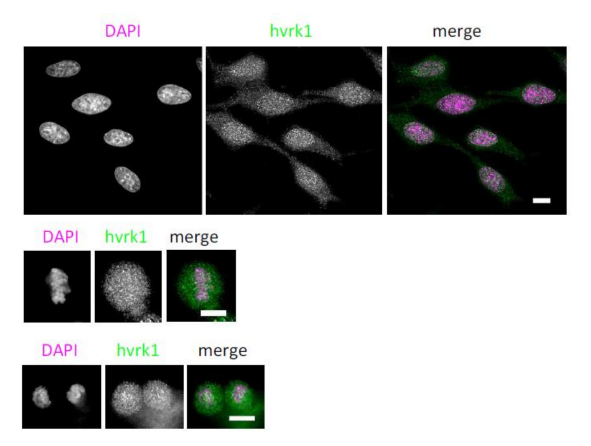


Fig.4. Immunoflorescence staining of VRK1 in HeLa cells, methanol fixed.

Top; Interphase cells were fixed with methanol and stained with the anti-human VRK1 antibody (hvrk1) at 1/100 dilution (center), DNA was stained with DAPI (left) and two images were merged (right; merge).

Center and bottom; Metaphase cells. At metaphase, VRK1 dots were solely detected in nuclei.



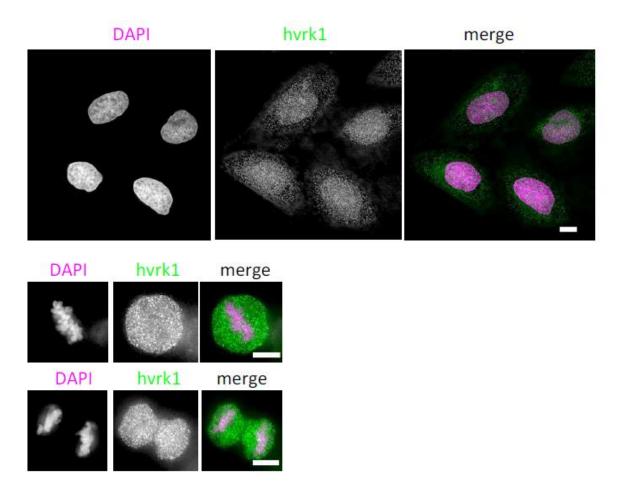


Fig. 5. Immunoflorescence staining of VRK1 in U-2 OS cells, formaldehyde fixed.

Top; Interphase cells were fixed with formaldehyde and stained with the anti-human VRK1 antibody (hvrk1) at 1/100 dilution (center), DNA was stained with DAPI (left) and two images were merged (right; merge).

Center and bottom; Metaphase cells. At metaphase, VRK1 dots were solely detected in nuclei.



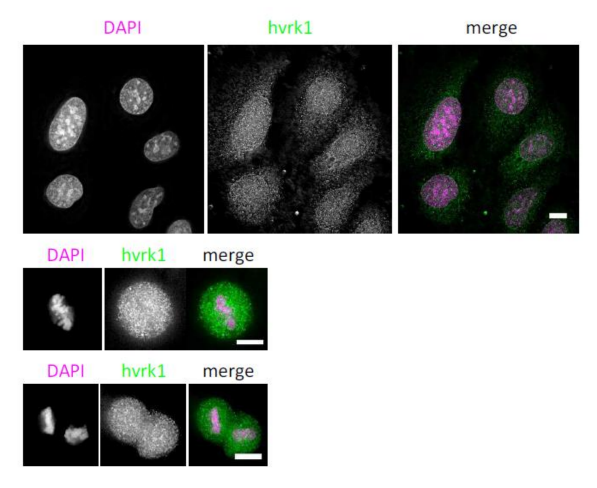


Fig. 6. Immunoflorescence staining of VRK1 in U-2 OS cells, methanol fixed.

Top; Interphase cells were fixed with methanol and stained with the anti-human VRK1 antibody (hvrk1) at 1/100 dilution (center), DNA was stained with DAPI (left) and two images were merged (right; merge).

Center and bottom; Metaphase cells. At metaphase, VRK1 dots were solely detected in nuclei.