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Datasheet for ABIN1381135 FGFR2 ELISA Kit



Overview

Quantity:	96 tests
Target:	FGFR2
Reactivity:	Human, Mouse, Rat
Method Type:	Cell ELISA
Application:	ELISA

Product Details

Sample Type:	Cell Culture Cells
Analytical Method:	Qualitative
Detection Method:	Colorimetric
Specificity:	The Anti-FGFR2 antibody is a rabbit polyclonal antibody. It was tested on Western Blots for
	specificity. A single protein band was detected. This protein band can be blocked by the
	synthesized immunogen peptide.
	The Anti-GAPDH antibody is a mouse monoclonal antibody. It was tested on Western Blots with
	the tissue lysates from human, mouse, and rat for specificity. A single protein band was
	detected from all three lysates.
Characteristics:	1) Cell Line: The cell line must express the target protein. This protocol can be used directly for
	adherent cells. For suspension cells and loosely attached cells, two steps are required: Coat the
	plates with 100 μ L of 10 μ g/mL Poly-L-Lysine to each well of the 96-well plate for 30 minutes at
	37°C before proceeding to Step 1 of Assay Protocol. Use 8% formaldehyde to fix the cells on
	Step 5 of Assay Protocol.

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	2) Cell Number and Sensitivity: The number of cells plated onto the 96-well plates depends on
	the expression level of FGFR2 protein in the cells, cell size, treatment conditions and incubation
	time. The cells used for testing should be around 75-90% confluent. Typically for HeLa cells,
	seed 30,000 cells per well overnight for treatment the following day. The FGFR2 Colorimetric
	Cell-Based ELISA Kit can detect FGFR2 expression in as little as 5,000 HeLa cells.
	3) Cell Treatment: The cells can be treated with inhibitors, activators, stimulators (ie. chemicals,
	proteins/peptides) or a combination of the substances listed above. The cells can be treated
	with UV and serum starvation to meet the needs of the end-user.
	4) Positive and Negative Controls: Mouse Anti-GAPDH antibody (included) should be used to
	detect the internal positive controls for normalization of OD values of the target protein. The
	negative controls are HRP-Conjugated Anti-Rabbit IgG antibody and HRP- Conjugated Anti-
	Mouse IgG antibody alone in different wells (without the primary antibodies). Both positive and
	negative controls should be performed in the same plate with the FGFR2 target experiments.
	5) Accuracy and Precision: Each condition should be performed in duplicate or in triplicate.
Components:	96-Well Cell Culture Clear-Bottom Microplate: 1 Plate
	TBS: 24 mL (10x), Clear
	Quenching Buffer: 24 mL (1x), Clear
	Blocking Buffer: 50 mL (1x), Clear
	Wash Buffer: 50 mL (10x), Clear
	Anti-FGFR2 antibody (Rabbit Polyclonal): 60 μl (100x), Purple
	Anti-GAPDH antibody (Mouse Monoclonal): 60 μl (100x), Green
	HRP-Conjugated Anti-Rabbit IgG antibody: 6 mL (1x), Glass
	HRP-Conjugated Anti-Mouse IgG antibody: 6 mL (1x), Glass
	Primary antibody Diluent: 12 mL (1x), Clear
	Ready-to-Use Substrate: 12 mL (1x), Brown
	Stop Solution: 12 mL (1x), Clear
	Crystal Violet Solution: 6 mL (1x), Glass
	SDS Solution: 24 mL (1x), Clear
	Adhesive Plate Seals: 4 Seals
Material not included:	The following materials and equipment are NOT provided in this kit but are necessary to
	successfully conduct the experiment:
	Microplate reader able to measure absorbance at 450 nm and/or 595 nm for Crystal Violet Cell

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Target Details

Target:	FGFR2
Alternative Name:	FGFR2 (FGFR2 Products)
Molecular Weight:	92025
OMIM:	176943
UniProt:	P21802
Pathways:	RTK Signaling, Fc-epsilon Receptor Signaling Pathway, EGFR Signaling Pathway, Neurotrophin Signaling Pathway, Regulation of Muscle Cell Differentiation, Skeletal Muscle Fiber Development, Growth Factor Binding

Application Details

Plate:	Uncoated
Protocol:	The Colorimetric Cell-Based ELISA Kit allows for the detection of various target proteins and the
	effects that certain stimulation conditions have on target protein expression in different cell
	lines. Qualitative determination of target protein concentration is achieved by an indirect ELISA
	format. In essence, the target protein is captured by target-specific primary (1st) antibodies
	while the HRP-conjugated secondary (2nd) antibodies bind the Fc region of the 1st antibody.
	Through this binding, the HRP enzyme conjugated to the 2nd antibody can catalyze a
	colorimetric reaction upon substrate addition. Due to the qualitative nature of the Cell-Based
	ELISA, multiple normalization methods are described:
	1) a monoclonal antibody specific for human GAPDH is included to serve as an internal positive
	control in normalizing the target absorbance values.
	2) Following the colorimetric measurement of HRP activity via substrate addition, the Crystal
	Violet whole-cell staining method is used to determine cell density. After staining, the results
	can be analyzed by normalizing the absorbance values to cell amounts, by which the plating
	difference can be adjusted.
	3) If a phosphorylated target is being detected, an antibody against the non- phosphorylated
	counterpart will be provided for normalization purposes. The absorbance values obtained for
	the non-phosphorylated target can be used to normalize the absorbance values for the

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FGFR2 Colorimetric Cell-Based ELISA

The FGFR2 Cell-Based ELISA Kit is a convenient, lysate-free, high throughput and sensitive assay kit that can monitor FGFR2 protein expression profile in cells. The kit can be used for measuring the relative amounts of FGFR2 in cultured cells as well as screening for the effects that various treatments, inhibitors (ie. siRNA or chemicals), or activators have on FGFR2.

Reagent Preparation:Note: Please remember to allow all solutions to warm up to room temperature prior to use.1x TBS: 1x TBS is used to wash cells seeded on the plate. 1x TBS can be prepared by adding 1volume of 10x TBS provided in the kit to 9 volumes of ddH20.

Fixing Solution: This solution is NOT provided. Fixing Solution is used to fix cells after cell culture. It is prepared by adding formaldehyde to 1x TBS with light mixing. The 4% formaldehyde is used for adherent cells and 8% formaldehyde is used for suspension cells and loosely attached cells.

Quenching Buffer: This solution is provided as ready-to-use. Quenching Buffer is used to inactivate the endogenous peroxidase activity of the seeded cells.

Blocking Buffer: This solution is provided as ready-to-use. Blocking Buffer is used to block additional binding sites in each well.

Wash Buffer: This buffer is provided as a 10x solution. 1x Wash Buffer can be prepared by adding 1 volume of 10x Wash Buffer provided in the kit to 9 volumes of ddH2O.

100x Anti-FGFR2 antibody: This antibody is a rabbit polyclonal antibody. This antibody was tested to be specific for the FGFR2 protein. The supplied antibody is a 100x solution. Make 1:100 dilutions in Primary antibody Diluent prior to use. The diluted primary antibody can be stored at 4°C for up to two weeks.

100x Anti-GAPDH antibody: This antibody is a mouse monoclonal antibody. This antibody was tested to be specific for GAPDH. The supplied antibody is a 100x solution. Make 1:100 dilutions in Primary antibody Diluent prior to use. The diluted primary antibody can be stored at 4°C for up to two weeks.

HRP-Conjugated Anti-Rabbit IgG antibody: This solution is provided as ready-to-use. HRP-Conjugated Anti-Rabbit IgG antibody is used as the secondary antibody to detect the targetbound, primary rabbit antibodies.

HRP-Conjugated Anti-Mouse IgG antibody: This solution is provided as ready-to-use. HRP-Conjugated Anti-Mouse IgG antibody is used as the secondary antibody to detect the targetbound, primary mouse antibodies.

Primary antibody Diluent: This solution is provided as ready-to-use. Use this solution to dilute the provided antibodies.

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	Ready-to-Use Substrate: This solution is provided as ready-to-use. Ready-to-Use Substrate
	must be warmed to room temperature before use. Keep away from light as this solution is light-
	sensitive.
	Stop Solution: This solution is provided as ready-to-use. Stop Solution must be handled with
	caution as it contains 2 N Sulfuric Acid (H2SO4) and is corrosive. Wear eye protection and
	gloves when handling.
	Crystal Violet Solution: This solution is provided as ready-to-use. Crystal Violet is an intense
	stain used to stain cell nuclei. Avoid contact with skin and clothing.
	SDS Solution: This solution is provided as ready-to-use. SDS is used to solubilize the Crystal
	Violet in preparation for cell staining. Store this solution at room temperature or warm up to
	room temperature if stored at 4°C.
Assay Procedure:	Note: Please read the whole manual before performing the experiment.
	1) Seed 200 μL of 20,000 adherent cells in culture medium in each well of a 96-well plate. The
	plates included in the kit are sterile and treated for cell culture. For suspension cells and loosely
	attached cells, coat the plates with 100 μL of 10 $\mu g/mL$ Poly-L-Lysine (not included) to each
	well of a 96-well plate for 30 minutes at 37°C prior to adding cells.
	2) Incubate the cells for overnight at 37°C, 5% CO2.
	3) Treat the cells as desired.
	4) Remove the cell culture medium and rinse with 200 μL of 1x TBS, twice.
	5) Fix the cells by incubating with 100 μL of Fixing Solution for 20 minutes at room
	temperature. The 4% formaldehyde is used for adherent cells and 8% formaldehyde is used for
	suspension cells and loosely attached cells. During the incubation, the plates should be sealed
	with Parafilm. Note: Fixing Solution is volatile. Wear appropriate personal protection equipment
	(mask, gloves and glasses) when using this chemical.
	6) Remove the Fixing Solution and wash the plate 3 times with 200 μL 1x Wash Buffer for five
	minutes each time with gentle shaking on the orbital shaker. The plate can be stored at 4°C for
	a week. Note: For all wash steps, tap the plate gently on absorbent papers to remove the
	solution completely.
	7) Add 100 μ L Quenching Buffer and incubate for 20 minutes at room temperature.
	8) Wash the plate 3 times with 1x Wash Buffer for 5 minutes at a time, with gentle shaking on
	the shaker.
	9) Add 200 μL of Blocking Buffer and incubate for 1 hour at room temperature. 10) Wash 3
	times with 200 μL of 1x Wash Buffer for 5 minutes at a time, with gentle shaking on the shaker.
	11) Add 50 μL of 1x primary antibodies (Anti-FGFR2 antibody and/or Anti-GAPDH antibody) to
	the corresponding wells, cover with Parafilm and incubate for 16 hours (overnight) at 4°C. If the

target expression is known to be high, incubate for 2 hours at room temperature with gentle shaking on the shaker.

12) Wash 3 times with 200 μL of 1x Wash Buffer for 5 minutes at a time, with gentle shaking on the shaker.

13) Add 50 µL of 1x secondary antibodies (HRP-Conjugated Anti- Rabbit IgG antibody and/or HRP-Conjugated Anti-Mouse IgG antibody) to corresponding wells and incubate for 1.5 hours at room temperature with gentle shaking on the shaker. Note: Add HRP-Conjugated Anti-Rabbit IgG antibody to the wells incubated with Anti-FGFR2 (rabbit, polyclonal) and add HRP-Conjugated Anti-Mouse IgG antibody to the wells incubated with Anti-GAPDH antibody (mouse, monoclonal).

14) Wash 3 times with 200 μ L of 1x Wash Buffer for 5 minutes at a time, with gentle shaking on the shaker.

15) Add 50 µL of Ready-to-Use Substrate to each well and incubate for 30 minutes at room temperature in the dark with gentle shaking on the shaker. Note: Ready-to-Use Substrate is a light-sensitive reagent. Keep away from light.

16) Add 50 µL of Stop Solution to each well and read OD at 450 nm immediately using the microplate reader. Optional: Crystal Violet Cell Staining Crystal Violet binds to cell nuclei and gives absorbance readings proportional to cell counts at 595 nm.

17) After finishing reading the absorbance at 450 nm, wash the plate twice with 200 μL of Wash Buffer and twice with 200 μL of 1x TBS for 5 minutes each. Tap the plates on paper towel to remove the excess liquid. Let plate air dry for 5 minutes at room temperature.
18) Add 50 μL of Crystal Violet Solution to each well, incubate for 30 minutes at room

temperature on the shaker. Note: Crystal Violet is an intense stain. Avoid contact with skin and clothing.

19) Flick the plate to remove Crystal Violet Solution, rinse the plate by filling the wells with running tap water, and wash the plate with 200 μ L of 1x TBS 3 times, 5 minutes each with gently shaking on the shaker.

20) Add 100 μ L of SDS Solution into each well and incubate on the shaker at room temperature for 1 hour.

21) Read absorbance at 595 nm with microplate reader. If absorbance is too high, the solubilized Crystal Violet Solution can be diluted 10 times with H2O on a separate 96-well plate.

Calculation of Results: GAPDH normalization: The OD450 values obtained for the target protein can be normalized using the OD450 values obtained for GAPDH. Crystal Violet Staining normalization: The measured OD450 readings can be normalized using the OD595 values via the proportion, OD450/OD595.

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