# Insulin Receptor [pY1334] ELISA Kit

Catalog Number KHR9161 (96 tests)

Pub. No. MAN0017201 Rev. 1.0 (30)

**CAUTION!** This kit contains materials with small quantities of sodium azide. Sodium azide reacts with lead and copper plumbing to form explosive metal azides. Upon disposal, flush drains with a large volume of water to prevent azide accumulation. Avoid ingestion and contact with eyes, skin and mucous membranes. In case of contact, rinse affected area with plenty of water. Observe all federal, state, and local regulations for disposal.

**Note:** For safety and biohazard guidelines, see the "Safety" appendix in the *ELISA Technical Guide* (Pub. no. MAN0006706). Read the Safety Data Sheets (SDSs) and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves.

## **Product description**

The Invitrogen<sup>™</sup> Insulin Receptor [pY1334] ELISA Kit is a solid-phase sandwich Enzyme-Linked Immunosorbent Assay (ELISA). This assay detects and quantifies the level of insulin receptor [pY1334] in cell lysates. The assay will recognize both natural and recombinant insulin receptor [pY1334].

Insulin receptor (IR), a cell surface receptor, binds insulin and mediates its action on target cells. The  $\beta$ -subunit (95 kDa) possesses a single transmembrane domain with tyrosine kinase activity. Phosphorylation of tyrosine 1328 and 1334 is required for the insulin-stimulated tyrosine dephosphorylation of pp125FAK and plays an inhibitory role in insulin-induced mitogenic signaling.

For normalizing IR content of the samples, an IR ( $\beta$ -subunit) ELISA kit independent of phosphorylation status is available (Cat.# KHR9111).

#### Contents and storage

Upon receipt, store the kit at 2°C to 8°C.

Contents	Cat. No. KHR9161 (96 tests)
IR [pY1334] Standard, lyophilized; contains 0.1% sodium azide. Refer to vial label for quantity and reconstitution volume	2 vials
Standard Diluent Buffer; contains 0.1% sodium azide; red dye <sup>[1]</sup>	25 mL
IR (ß-subunit) Antibody Coated Plate, 96-well strip-well plate	1 plate
IR [pY1334] Detection Antibody; contains 0.1% sodium azide; blue dye <sup>[1]</sup>	11 mL
Anti-Rabbit IgG HRP (100X); contains 3.3 mM thymol	0.125 mL
HRP Diluent; contains 3.3 mM thymol; yellow dye <sup>[1]</sup>	25 mL
Wash Buffer Concentrate (25X)	100 mL
Stabilized Chromogen, Tetramethylbenzidine (TMB)	25 mL
Stop Solution	25 mL
Plate Covers, adhesive strips	3

[1] In order to help our customers avoid any mistakes in pipetting the ELISAs, we provide colored Standard Diluent Buffer, Detection Antibody, and HRP Diluent to help monitor the addition of solution to the reaction well. This does not in any way interfere with the test results.

## **Required materials not supplied**

- Distilled or deionized water
- Microtiter plate reader with software capable of measurement at or near 450 nm
- Plate washer–automated or manual (squirt bottle, manifold dispenser, or equivalent)
- Calibrated adjustable precision pipettes and glass or plastic tubes for diluting solutions; beakers, flask and cylinders for preparation of reagents
- Cell Extraction Buffer (Cat. No. FNN0011, or see "Prepare Cell Extraction Buffer")

## Before you begin

**IMPORTANT!** Reagents are lot-specific. Do not mix or interchange different reagent lots from various kit lots.

- Review the **Procedural guidelines** and **Plate washing directions** in the *ELISA Technical Guide* available at **thermofisher.com**.
- Allow reagents to reach room temperature before use. Mix to redissolve any precipitated salts.

## Prepare 1X Wash Buffer

- 1. Dilute 16 mL of Wash Buffer Concentrate (25X) with 384 mL of deionized or distilled water. Label as 1X Wash Buffer.
- 2. Store the concentrate and 1X Wash Buffer in the refrigerator. Use the diluted buffer within 14 days.

## **Prepare Cell Extraction Buffer**

**Note:** See the *ELISA Technical Guide* for detailed information on preparing Cell Extraction Buffer.

1. Prepare Cell Extraction Buffer.

Cell Extraction Buffer consists of 10 mM Tris (pH 7.4), 100 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1 mM NaF, 20 mM Na<sub>4</sub>P<sub>2</sub>O<sub>7</sub>, 2 mM Na<sub>3</sub>VO<sub>4</sub>, 1% Triton<sup>™</sup> X-100, 10% glycerol, 0.1% SDS, and 0.5% deoxycholate.

 Immediately before use, add 1 mM PMSF (0.3 M stock in DMSO) and 100 μL of protease inhibitor cocktail (e.g., Sigma Cat. No. P-2714) per 1 mL of Cell Extraction Buffer.



## Prepare cell lysate

- 1. Collect cells by centrifugation (non-adherent cells) or scraping from culture flasks (adherent cells), then wash cells twice with cold PBS.
- 2. Remove and discard the supernatant and collect the cell pellet. The pellet can be stored at -80°C and lysed at a later date if desired.
- 3. Lyse the cell pellet in Cell Extraction Buffer for 30 minutes, on ice. Vortex at 10-minute intervals.
- Note: The volume of Cell Extraction Buffer used depends on the number of cells in the cell pellet, and expression levels of insulin receptor [pY1334].
- 4. Transfer the lysate into microcentrifuge tubes and centrifuge at 13,000 rpm for 10 minutes at 4°C.
- 5. Transfer the supernatant into clean microcentrifuge tubes. Samples can be stored at -80°C (avoid multiple freeze-thaw cycles).

## **Pre-dilute samples**

Sample concentrations should be within the range of the standard curve. Because conditions may vary, each investigator should determine the optimal dilution for each application.

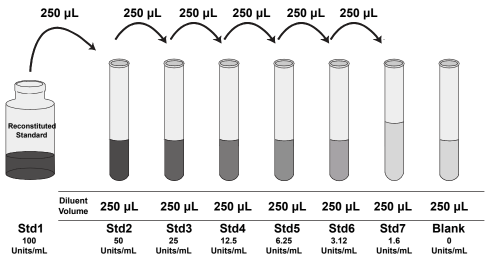
- Perform sample dilutions with Standard Diluent Buffer.
- Dilute samples prepared in Cell Extraction Buffer 1:10 or greater in Standard Diluent Buffer (e.g., 10 µL sample into 90 µL buffer). This dilution is necessary to reduce the matrix effect of the Cell Extraction Buffer. SDS concentration should be less than 0.01% before adding to the plate. While a 1:10 sample dilution has been found to be satisfactory, higher dilutions such as 1:25 or 1:50 may be optimal.

## **Dilute standards**

Note: Use glass or plastic tubes for diluting standards.

**Note:** This IR [pY1334] Standard was prepared from purified, phosphorylated  $\beta$ -subunit of IR expressed in Sf9 cells. One unit of standard is equivalent to the amount of IR [pY1334] autophosphorylated from 60 pg of IR ( $\beta$ -subunit) protein.

- 1. Reconstitute IR [pY1334] Standard to 100 Units/mL with Standard Dilution Buffer. Refer to the standard vial label for instructions. Swirl or mix gently and allow the contents to sit for 10 minutes to ensure complete reconstitution. Label as 100 Units/mL insulin receptor [pY1334]. Use the standard within 1 hour of reconstitution.
- 2. Add 250 µL Standard Diluent Buffer to each of 7 tubes labeled as follows: 50, 25, 12.5, 6.25, 3.12, 1.6, and 0 Units/mL insulin receptor [pY1334].
- 3. Make serial dilutions of the standard as shown in the following dilution diagram. Mix thoroughly between steps.
- 4. Remaining reconstituted standard should be discarded or frozen in aliquots at -80°C for further use. Standard can be frozen and thawed one time only without loss of immunoreactivity.



## Prepare 1X Anti-Rabbit IgG HRP solution

Note: Prepare 1X Anti-Rabbit IgG HRP solution within 15 minutes of usage.

- The Anti-Rabbit IgG HRP (100X) is in 50% glycerol, which is viscous. To ensure accurate dilution:
- For each 8-well strip used in the assay, pipet 10 μL Anti-Rabbit IgG HRP (100X) solution, wipe the pipette tip with clean absorbent paper to remove any excess solution, and dispense the solution into a tube containing 1 mL of HRP Diluent. Mix thoroughly.
- 2. Return the unused Anti-Rabbit IgG HRP (100X) solution to the refrigerator.

# Perform ELISA (Total assay time: 4 hours)

**IMPORTANT!** Perform a standard curve with each assay.

- Allow all components to reach room temperature before use. Mix all liquid reagents prior to use.
- Determine the number of 8-well strips required for the assay. Insert the strips in the frames for use. Re-bag any unused strips and frames, and store at 2°C to 8°C for future use.

Y Cap anti	ture 🔨 Antigen 🩏 Detector body 🔪 Antigen	HRP Secondary antibody
1	Bind antigen	a. Add 100 μL of standards, controls, or samples (see "Pre-dilute samples" on page 2) to the appropriate wells. Leave the wells for chromogen blanks empty.
	N.N. N.L.	b. Cover the plate with a plate cover and incubate 2 hours at room temperature.
		c. Thoroughly aspirate the solution and wash wells 4 times with 1X Wash Buffer.
2	Add detector antibody	a. Add 100 $\mu$ L of IR [pY1334] Detection Antibody solution into each well except the chromogen blanks.
~		<b>b</b> . Cover the plate with a plate cover and incubate 1 hour at room temperature.
	×	<b>c.</b> Thoroughly aspirate the solution and wash wells 4 times with 1X Wash Buffer.
3	Add IgG HRP	a. Add 100 µL Anti-Rabbit IgG HRP solution into each well except the chromogen blanks.
5		b. Cover the plate with plate cover and incubate for 30 minutes at room temperature.
	y sk	<b>c</b> . Thoroughly aspirate the solution and wash wells 4 times with 1X Wash Buffer.
1.	Add Stabilized Chromogen	a. Add 100 µL Stabilized Chromogen to each well. The substrate solution begins to turn blue.
		<b>b.</b> Incubate for 30 minutes at room temperature in the dark.
	A A A	Note: TMB should not touch aluminum foil or other metals.
5	Add Stop Solution	Add 100 $\mu$ L Stop Solution to each well. Tap the side of the plate to mix. The solution in the wells changes
5	<b>X</b>	from blue to yellow.

## Read the plate and generate the standard curve

- 1. Read the absorbance at 450 nm. Read the plate within 2 hours after adding the Stop Solution.
- 2. Use curve-fitting software to generate the standard curve. A 4 parameter algorithm provides the best standard curve fit. Optimally, the background absorbance may be subtracted from all data points, including standards, unknowns and controls, prior to plotting.
- 3. Read the concentrations for unknown samples and controls from the standard curve. Multiply value(s) obtained for sample(s) by the appropriate factor to correct for the sample dilution.

**Note:** Dilute samples producing signals greater than the upper limit of the standard curve in Standard Diluent Buffer and reanalyze. Multiply the concentration by the appropriate dilution factor.

## Performance characteristics

#### Standard curve example

The following data were obtained for the various standards over the range of 0 to 100 Units/mL insulin receptor [pY1334].

Standard Insulin Receptor [pY1334] (Units/mL)	Optical Density (450 nm)
100	2.96
50	1.60
25	0.85
12.5	0.43
6.25	0.25
3.12	0.19
1.6	0.15
0	0.10

#### Recovery

To evaluate recovery, insulin receptor [pY1334] Standard was spiked at 3 different concentrations into 10% Cell Extraction Buffer. The average percent recovery was 125%.

#### Inter-assay precision

Samples were assayed 48 times in multiple assays to determine precision between assays.

Parameters	Sample 1	Sample 2	Sample 3
Mean (U/mL)	49.72	23.23	9.21
Standard Deviation	2.48	1.54	0.72
% Coefficient of Variation	4.98	6.64	7.77

#### Intra-assay precision

Samples of known insulin receptor [pY1334] concentrations were assayed in replicates of 16 to determine precision within an assay.

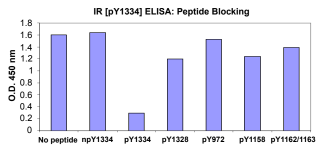
Parameters	Sample 1	Sample 2	Sample 3
Mean (U/mL)	50.02	24.25	9.17
Standard Deviation	2.07	0.93	0.48
% Coefficient of Variation	4.14	3.83	5.19

#### **Cross-reactivity**

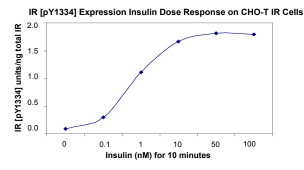
Although this ELISA kit was developed using human IR transfected cells, cross-reactivity with mouse and rat insulin receptor is documented.

#### Specificity

Insulin Receptor [pY1334] ELISA Kit specificity was confirmed by peptide competition. Data show only phosphopeptide containing the phosphorylated tyrosine blocks the ELISA signal. Same sequence with a non-phosphorylated tyrosine at position 1334 was not blocked.



IR [pY1334] phosphorylation in CHO-T IR cells is dependent on the level of insulin stimulation. Cells (~90% confluent) were treated with insulin at varying concentrations (0 to 100 nM) for 10 minutes, lysed, and measured in parallel for IR ( $\beta$ -subunit) (Cat.# KHR9111) and IR [pY1334] content. IR [pY1334] levels were normalized to total IR ( $\beta$ -subunit).



## Parallelism

Natural insulin receptor [pY1334] from 50 nM insulin-treated CHO-T IR cells was serially diluted in Standard Diluent Buffer. The optical density of each dilution was plotted against the insulin receptor [pY1334] standard curve. The standard accurately reflects the full insulin receptor [pY1334] content in samples.

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Product	t label explana	tion of s	ymbols and wa	rnings							
REF	Catalog Number	LOT	Batch code	1	Temperature limitation	Use by	Manufacturer	i	Consult instructions for use	$\triangle$	Caution, consult accompanying documents

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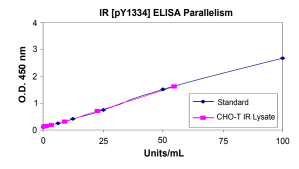
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#### Sensitivity

The analytical sensitivity of insulin receptor [pY1334] is <1 Unit/mL as determined by adding two standard deviations to the mean O.D. obtained from 30 assays of the zero standard. This corresponds to the insulin receptor [pY1334] extracted from 3,000 CHO-T cells treated with 50 nM insulin for 10 minutes. ELISA sensitivity is ~4-fold greater than Western blotting when tested against known quantities of insulin receptor [pY1334].

IR [pY1334] (95 kDa)								
ELISA: O.D. 450 nm	0.21	0.23	0.26	0.28	0.37	0.62	1.3	2.2
CHO-T IR lysate (µq/test)	0	0.313	0.625	1.25	2.5	5	10	20

#### Linearity of dilution

CHO-T IR cells were treated with 50 nM insulin for 10 minutes then harvested and lysed with Cell Extraction Buffer. This lysate was diluted in Standard Diluent Buffer over the assay range and measured for insulin receptor [pY1334]. Linear regression analysis of sample values vs. expected concentrations yielded a correlation coef. of 0.99.

Dilution	Measured (U/mL)	Expected				
Ditution	Measureu (O/IIIL)	(U/mL)	%			
Neat	45.21	45.22	100.00			
1/2	21.02	22.61	92.94			
1/4	9.71	11.30	85.92			
1/8	4.50	5.65	79.53			



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