Direct Detection of His-tagged Proteins Using Nickel-NTA Conjugates

For Products:

KPL HisDetector [™] Western Blot Kits	Catalog No.
KPL HisDetector [™] Western Blot Kit, HRP	5810-0023
Colorimetric	(24-00-01)
HisDetector [™] Western Blot Kit, HRP	5810-0024
Chemiluminescent	(24-00-02)
HisDetector [™] Western Blot Kit, AP Colorimetric	5810-0025 (25-00-01)

KPL HisDetector [™] Nickel Conjugates	Catalog No.
HisDetector [™] Nickel-HRP	5820-0001 (24-01-01)
HisDetector [™] Nickel-AP	5820-0003 (25-01-01)

Store KPL HisDetector[™] Nickel-HRP at -20°C Immediately Upon Arrival



TABLE OF CONTENTS

Section	Page
Product Contents	3
Storage and Stability	4
KPL HisDetector [™] Product Description Using His-tagged Proteins Nickel Detection of His-tagged Proteins	5
Before You Begin Safety and Handling Required Supplies and Equipment Important Product Information	7
Reagent Preparation	8
Western Blot and Dot Blot Protocols Western Blot Protocol at a Glance Sample Preparation and Gel Electrophoresis HRP Western Blot Detection AP Western Blot Detection Dot Blot Detection Stripping and Re-probing Blots	9
ELISA Protocols ELISA Protocol at a Glance ELISA Plate Coating HRP ELISA Detection AP ELISA Detection	14
Troubleshooting Guide KPL HisDetector Western Blot Protocols KPL HisDetector ELISA Protocols	18
References	22
Related Products	22
Limited Use Label and Disclaimers	23

PRODUCT CONTENTS

WESTERN BLOT KITS

5810-0023 (24-00-01) HisDetector[™] Western Blot Kit, HRP Colorimetric

Size: 40 Blots

Kit Component	Part Number	Volume	Quantity
KPL HisDetector [™] Nickel-HRP	5820-0002 (24-01-02)	0.1 mL	4
KPL BSA Powder	5810-0027 (51-61-00)	10 grams	1
KPL 20X TBST	5810-0026 (50-63-16)	100 mL	2
TMB 1-Component Membrane Substrate	5120-0055 (50-77-02)	100 mL	4

5810-0024 (24-00-02) KPL HisDetector[™] Western Blot Kit, HRP Chemiluminescent

Size: 40 Blots			
Kit Component	Part Number	Volume	Quantity
KPL HisDetector [™] Nickel-HRP	5820-0002 (24-01-02)	0.1 mL	1
KPL BSA Powder	5810-0027 (51-61-00)	10 grams	1
KPL 20X TBST	5810-0026 (50-63-16)	100 mL	2
KPL LumiGLO Substrate, Solution A	5430-0027 (50-59-00)	120 mL	1
KPL LumiGLO Substrate, Solution B	5430-0030 (50-60-00)	120 mL	1

5810-0025 (25-00-01) HisDetector[™] Western Blot Kit, AP Colorimetric

Kit Component	Part Number	Volume	Quantity
KPL HisDetector [™] Nickel-AP	5820-0004 (25-01-02)	1 mL	1
KPL 5X Detector Block	5440-0003 (71-83-01)	120 mL	2
KPL 20X TBST	5810-0026 (50-63-16)	100 mL	2
KPL BCIP/NBT 1-Component Membrane Substrate	5420-0034 (50-81-08)	100 mL	4

KPL HisDetector Western Blot Kits provide sufficient reagents to test approximately 4000 cm² of membrane (forty 10 cm x 10 cm blots) when using recommended volumes. If desired, increased working volumes may be used; however, additional reagents will be necessary. See Related Products on page 22 for information on additional reagents.

PRODUCT CONTENTS (cont.)

Ni-NTA CONJUGATES

5820-0001 (24-01-01) KPL HisDetector[™] Nickel-HRP

Size: 0.1 mL			
Component	Part Number	Volume	Quantity
KPL HisDetector [™] Nickel-HRP	5820-0002 (24-01-02)	0.1 mL	1

5820-0003 (25-01-01) KPL HisDetector[™] Nickel-AP

Size: 1 mL

Component	Part Number	Volume	Quantity
KPL HisDetector [™] Nickel-AP	5820-0004 (25-01-02)	1.0 mL	1
KPL Detector [™] Block (5X)	5440-0003 (71-83-01)	120 mL	1

STORAGE AND STABILITY

Storage

KPL HisDetector[™] Nickel-AP products are shipped at ambient temperature. Upon receipt, store KPL HisDetector[™] Nickel-AP kits at 2-8°C.

KPL HisDetector[™] Nickel-HRP products are shipped with cold pack. Upon receipt, store KPL HisDetector[™] Nickel-HRP (24-01-02) at -20°C and other kit components at 2-8°C.

Stability

KPL HisDetector[™] Western Blot Kits and Conjugates are stable for a minimum of one year from date of receipt when stored at recommended temperatures.

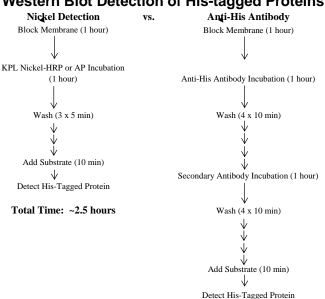
KPL HisDetector[™] PRODUCT DESCRIPTION

Using His-Tagged Proteins

His-tagged fusion is commonly employed in protein analysis to facilitate the production, purification, and detection of a protein of interest. The first step in this process is typically inserting a DNA sequence coding for four to ten consecutive histidines into the coding region flanking the gene of the protein sequence being expressed. The protein containing the His-tag fusion can then be expressed in large quantities and easily purified over immobilized metal affinity chromatography (IMAC) resins. In addition, the His-tag can be further used in downstream applications to determine level of protein expression, protein function, and proteinprotein interactions.

Nickel Detection of His-Tagged Proteins

His-tagged proteins have a high affinity for nickel (Ni²⁺) ions. Nickel-NTA (nitrilotriacetic acid) conjugated to horseradish peroxidase (HRP) or alkaline phosphatase (AP) will therefore bind directly to the His-tag, eliminating the need for primary and secondary antibodies. This results in assay simplification and reduced detection time. Direct detection with nickel-NTA also offers excellent sensitivity and low background in applications such as Western blotting. Nickel-NTA conjugates can detect proteins with a His-tag at either their N- or C-terminus.



Western Blot Detection of His-tagged Proteins

Total Time: ~4.5 hours

L-1004175-01 December 2017

KPL HisDetector[™] PRODUCT DESCRIPTION (cont.)

KPL HisDetector Nickel Conjugates

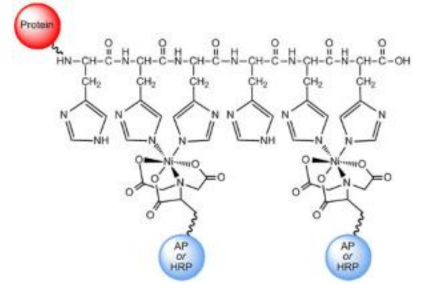
KPL HisDetector Nickel-HRP and KPL Nickel-AP conjugates are tools for detection of His-tagged recombinant proteins. These products are made by covalently conjugating nickel-NTA to the reporter enzymes horseradish peroxidase (HRP) and alkaline phosphatase (AP), respectively. Each conjugate is manufactured to provide high sensitivity and low background in Western blotting applications using both chemiluminescent and colorimetric substrates. KPL HisDetector nickel conjugates may also be used in dot blotting, ELISA and IHC applications.

KPL HisDetector Western Blot Kits

KPL HisDetector Western Blot Kits provide convenience and flexibility for researchers by offering KPL HisDetector nickel conjugates, assay reagents (blocker, wash solutions), and substrates within a fully optimized kit system. HRP Kits are offered for colorimetric and chemiluminescent detection. The KPL AP HisDetector[™] Western Blot Kit offers colorimetric detection.

His-tagged Protein Interaction with Ni-NTA Conjugated to HRP/AP:

Nickel ions (Ni^{2+}) can form six electron coordination bonds. NTA binds to four of these bonds. Two ligand binding sites are then available to readily coordinate with a 6X His-tagged protein as shown in the schematic below.



BEFORE YOU BEGIN

Safety and Handling

- Read SDS and all instructions thoroughly before using product.
- Wear appropriate personal protective equipment when handling all reagents.
- Caution should be used when preparing or handling polyacrylamide gels; monomeric acrylamide is a neurotoxin.

Required Supplies and Equipment

- Nitrocellulose, PVDF or Nylon membrane
- Polyacrylamide gels
- Electrophoresis equipment
- ß-mercaptoethanol or DTT
- Incubation trays or tubes for reagent incubation
- Platform shaker or rocker
- Coomassie blue for gel staining (optional)
- Protein stain such as Ponceau S (optional)
- Protein standards (optional)
- X-ray film (double emulsion such as Kodak BioMax[®] Light) or a Chemiluminescent Imaging System (for chemiluminescent detection only)
- Film developing chemicals and equipment (for chemiluminescent detection only)

Important Product Information

- His-tag segments of proteins may fold in a manner that reduces interaction between the His-tag and the nickel-NTA conjugate. Denaturing the sample by reducing all available disulfide bonds may enhance exposure of buried histidine residues and increase the probability of His-tag interaction with nickel-NTA. See Sample Preparation on page 10 for information on reducing samples.
- Use of certain buffers or reagents may inhibit interaction of HisDetector Nickel conjugates with the His-tagged segment of the protein. The most notable examples are:
 - Imidazole will bind to various divalent metal cations including nickel. It is commonly used to elute His-tagged proteins when bound to nickel resins.
 - EDTA and other divalent metal chelators will strip the nickel from the reporter enzyme. All divalent metal chelators should be avoided in nickel detection systems.

REAGENT PREPARATION

• KPL 1% BSA Blocking Solution

Dissolve KPL BSA Powder in KPL 1X TBST Wash Solution to a final concentration of 1% (w/v), i.e., 1 gram of KPL BSA Powder in 100 mL KPL 1X TBST Wash Solution. Freshly prepare solution for each use and discard any unused portion. Alternative blocking solutions may significantly reduce sensitivity, especially if they contain milk-based or phosphorylated proteins.

• KPL 1X Detector Block Solution

Dilute KPL Detector Block Solution (5X) 1/5 v/v in reagent quality water. Freshly prepare solution for each use and discard any unused portion. **Do not use an alternative blocking solution in applications using KPL Nickel-AP.**

• KPL 1X TBST Wash Solution

Dilute KPL 20X TBST Wash Solution Concentrate 1/20 with reagent quality water (*i.e.*, 10 mL Wash Solution Concentration + 190 mL H₂O). Store at room temperature or at 2 - 8°C. **Do not use wash solution with EDTA or imidazole.**

KPL LumiGLO Chemiluminescent Substrate

Mix equal parts of KPL LumiGLO Solution A and KPL LumiGLO Solution B (*i.e.*, 5 mL Solution A + 5 mL Solution B). The KPL LumiGLO mixture is stable for up to 24 hours at $2-8^{\circ}$ C.

WESTERN BLOT PROTOCOLS

HisDetector[™] Western Blot Protocol at a Glance

Protein Gel



- 1. Polyacrylamide Gel Electrophoresis
- 2. Transfer Protein onto Membrane

3. Block Membrane, 1 hour

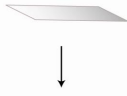
4. Incubate HisDetector Nickel Conjugate,

5. Wash Membrane, 3 x 5 minutes

6. Incubate Substrate, 1 - 10 minutes

1 hour

Western blot



Incubations





His-tagged protein results



 Visualize Bands (Colorimetric) or Expose to Film (Chemiluminescent),
 2 - 10 minutes





SAMPLE PREPARATION AND GEL ELECTROPHORESIS

 Prepare protein extract for electrophoresis. Add 5% βmercaptoethanol or 1% DTT (final concentration) to reduce protein samples. Incubate protein samples at 90-100°C for 3 minutes prior to loading on gel. The recommended amount of sample to load in each lane is 1-10 µg of crude lysate or 500 pg - 1 µg of purified protein. Optimization may be required depending on the expression of the His-tagged protein and substrate used.

Avoid overloading protein samples as it will adversely affect membrane detection results.

Recommended Control: It is advisable to run control samples containing protein extract from the expression strain that does not contain the recombinant His-tagged protein.

- 2. Electrophorese until tracking dye approaches the bottom of the gel. Conditions for electrophoresis will vary depending on the type of gel and the molecular weight of the protein off interest. Check the apparatus manufacturer for recommendations.
- Transfer proteins on to nitrocellulose, PVDF, or nylon membrane. Run transfer according to equipment manufacturer's instructions. KPL HisDetector[™] products have been optimized with nitrocellulose.

Optimal transfer time should be determined experimentally depending on the size and abundance of the protein of interest.

4. a). Optional: Stain the gel post-transfer with Coomassie[™] Blue to determine transfer efficiency.

b). Optional: Stain proteins with Ponceau-S for 10 minutes at room temperature with shaking. Use a sufficient volume of stain to cover the membrane. Remove membrane from stain and rinse with reagent quality water to remove excess stain. Protein bands will appear as background diminishes. Do not continue to rinse or specific protein staining will diminish.

The presence of proteins on the gel indicates sub-optimal transfer. Pre-stained protein markers can be used to reliably monitor the efficiency of transfer.

Ponceau-S for total protein stain is compatible with downstream detection, since it can be removed by washing the membrane in reagent quality water.

5. Proceed to detection on pages 11 and 12.

HRP WESTERN BLOT DETECTION

Volumes indicated are for a $10 \times 10 \text{ cm}^2$ blot and may be adjusted for other sizes or multiple blots. Perform all steps at room temperature.

- Block the membrane by completely immersing in 20 mL of freshly prepared KPL 1% BSA Block Solution for 1 hour at room temperature with gentle agitation or overnight at 2-8°C without agitation. Alternative blocking solutions may significantly reduce sensitivity, especially if they contain milk-based or phosphorylated proteins.
- 2. Add KPL HisDetector[™] Nickel-HRP conjugate.

Colorimetric Assay: Dilute KPL HisDetector Nickel-HRP 1/2,000-1/10,000 directly into the 20 mL of KPL 1% BSA Block Solution used to block the membrane. Incubate for 1 hour at room temperature with gentle agitation.

Chemiluminescent Assay: Dilute KPL HisDetector Nickel-HRP 1/10,000-1/40,000 directly into the 20 mL of KPL 1% BSA Block Solution used to block the membrane. Incubate for 1 hour at room temperature with gentle agitation.

Recommended dilutions are provided as starting points. It may be necessary to titrate the concentration of Nickel-HRP to receive optimal results with your system.

- 3. Wash the membrane by immersing in KPL 1X TBST, 3 times for 5 minutes each with gentle rocking.
- Colorimetric Detection: Add 10 mL KPL TMB Membrane Substrate directly to the membrane and allow to develop for 5-15 minutes. Rinse the membrane for 10-30 seconds in reagent quality water to stop the reaction. Allow the membrane to air dry.

A digital photograph is recommended if a permanent record is required.

Chemiluminescent Detection: Apply 5 mL KPL LumiGLO[®] working solution directly to the membrane for 1 minute. Remove the membrane from substrate and touch the corner to a piece of filter paper to remove excess liquid. Seal the membrane in clear plastic and expose to X-ray film for 2 - 10 minutes. An initial exposure of 2 minutes is recommended.

AP WESTERN BLOT DETECTION

Volumes indicated are for a $10 \times 10 \text{ cm}^2$ blot and may be adjusted for other sizes or multiple blots. Perform all steps at room temperature.

- Block the membrane by completely immersing in 20 mL KPL 1X Detector Block Solution for 1 hour at room temperature with gentle agitation or overnight at 2-8°C without agitation. Do not use alternative blocking solutions.
- 2. Add HisDetector[™] Nickel-AP conjugate.

Colorimetric Assay: Dilute KPL HisDetector Nickel-AP 1/1,000-1/2,000 directly into the 20 mL of KPL 1X Detector Block Solution used to block the membrane. Incubate for one hour at room temperature with gentle agitation.

Chemiluminescent Assay: Dilute KPL HisDetector Nickel-AP 1/10,000 to 1/40,000 directly in the 20 mL KPL 1X Detector Block Solution used to block the membrane. Incubate for one hour at room temperature with gentle agitation.

Recommended dilutions are provided as starting points. It may be necessary to titrate the concentration of KPL Nickel-AP to receive optimal results with your system.

- 3. Wash the membrane by immersing in KPL 1X TBST, 3 times for 5 minutes each with gentle rocking.
- 4. **Colorimetric Detection:** Incubate membrane in 10 mL KPL BCIP/NBT and allow to develop for 5-15 minutes. Rinse the membrane for 10-30 seconds in reagent quality water to stop the reaction. Allow the membrane to air dry.

A digital photograph is recommended if a permanent record is required.

Chemiluminescent Detection: Apply 5 mL AP Chemiluminescent Substrate directly to the membrane and incubate for 1 minute. Remove the membrane from substrate and touch the corner to a piece of filter paper to remove excess liquid. Seal the membrane in clear plastic and expose to X-ray film. For most applications, 1-10 minutes exposure is sufficient.

SeraCare recommends KPL PhosphaGLO AP Substrate for chemiluminescent detection of AP conjugates in Western blotting.

DOT BLOT DETECTION

The "Dot Blot" is a useful semi-quantitative technique for detecting and characterizing proteins. It is similar to Western blotting. However, the protein samples are directly spotted onto the membrane, instead of being transferred from a gel after electrophoresis. The Dot Blot can also be used to troubleshoot or optimize your Western blot. The sensitivity of detection is comparable between a Dot Blot and a Western Blot. However, the sensitivity tends to be slightly higher in Western blots. Dot Blots are especially useful when titrating the amount of conjugate or comparing various block solutions.

- Using a pencil, draw a grid to indicate the region you are going to blot. For 1 µL spots, draw squares at least ¼"x¼". We recommend using nitrocellulose membranes for dot blots.
- 2. For each protein, spot 1 μ L of the serially diluted protein samples onto the membrane. Use TBS as the diluent. In general, eight two-fold dilutions from 1 μ g/ μ L will yield good results.

Recommended Controls: It is advisable to run a positive control containing the 6xHis Protein Ladder (Qiagen), where 1 μ L corresponds to 20-25 ng of protein. The recommended negative control is protein extract that does not contain the recombinant Histagged protein.

- 3. Allow the membranes to air dry at least 2 hours before blocking.
- Refer to detection on pages 11 and 12. Note that dot blots do not typically require an entire 10 x 10 cm² membrane, so the reagent volumes can be reduced.

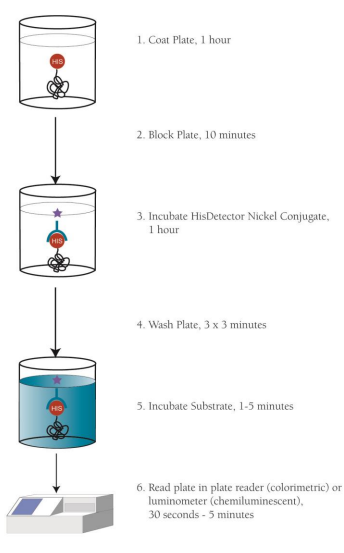
STRIPPING AND REPROBING BLOTS

It is possible to strip a membrane that has been probed with either KPL Nickel-HRP or KPL Nickel-AP and detected with KPL LumiGLO or KPL PhosphaGLO, respectively. It may be re-probed with an alternate detection method if the membrane is not allowed to dry prior to stripping. Stripping the membrane may be done several times, although a gradual increase in background and/or a decrease in sensitivity may occur.

- 1. Immerse blot in either 20 mL of 6M urea or 100 mM imidazole for 30 minutes with gentle shaking.
- 2. Carefully rinse blot in reagent quality water for 2 minutes.
- Block membrane by immersing in 20 mL of appropriate blocking solution for 1 hour at room temperature with gentle rocking or stationary at 2-8°C overnight.
- 4. Detect membrane as described on pages 11 and 12.

ELISA PROTOCOL

HisDetector[™] ELISA at a Glance Total time: 2.5 hours





ELISA PLATE COATING

Protocol describes coating purified His-tagged protein for ELISA (standard 96-well plate). Perform all steps at room temperature.

1. Dilute the His-tagged protein to 10 μ g/mL in PBS. Add 100 μ L per well and incubate for one hour.

The recommendations stated here offer a starting point. Optimal protein concentration must be pre-determined prior to coating. The amount of protein coated onto the plate will depend on the expression of the target protein.

SeraCare recommends the use of high binding microwell plates (i.e. Nunc MaxiSorp[™] or equivalent).

 Add 200 µL of blocking solution to each well and incubate for at least ten minutes.

SeraCare recommends: 1% BSA in 1X TBST for blocking ELISA plates.

- 3. Remove liquid from wells.
- 4. Add 200 µL/well of 2% sucrose and incubate for 5 minutes.
- 5. Remove liquid from wells and dry for at least 2 hours. Detect histagged proteins as described on page 16 or 17.

Coated plates can be stored dessicated for several months.

HRP ELISA DETECTION

Protocol is for a standard 96-well plate. Perform all steps at room temperature.

- Dilute KPL HisDetector[™] Nickel-HRP 1/1,000 to 1/10,000 in 1% BSA in 1X TBST. Add 100 µL diluted KPL Nickel-HRP per well. Incubate for 30 minutes.
- Wash plate by adding 300 µL KPL 1X TBST Wash Solution per well. Let plate stand for 3 minutes with wash solution in the wells. Remove wash solution from wells and repeat two additional times.

Other wash solutions may be used, however, the solution must not contain chelators of divalent metal cations, i.e., no EDTA.

 Add 100 µL/well of KPL ABTS Substrate for detection. Follow the product instructions for incubation time and necessity of stop solution.

> Other substrates are available for HRP detection in ELISA. SeraCare offers KPL TMB 2-Component, KPL SureBlue, or KPL SureBlue Reserve for colorimetric ELISA detection. For chemiluminescent detection, KPL LumiGLO or KPL LumiGLO Reserve are available.

> See RELATED PRODUCTS on page 22 for more information on HRP substrates for ELISA.

- Stop reaction by adding 100 µl of KPL ABTS Peroxidase Stop Solution (See RELATED PRODUCTS) or 1% Sodium Dodecyl Sulfate (SDS) to each well. ABTS substrate will remain blue-green after addition of stop solution.
- Read plate using a plate reader at the appropriate wavelength (405-410 nm for KPL ABTS) or a luminometer (chemiluminescent detection).

AP ELISA DETECTION

Protocol is for a standard 96-well plate. Perform all steps at room temperature.

- Dilute KPL HisDetector[™] Nickel-AP 1/1,000 to 1/10,000 in KPL 1X Detector Block. Add 100 µL diluted Nickel-AP per well. Incubate for 30 minutes.
- Wash plate by adding 300 µL KPL 1X TBST Wash Solution per well. Let plate stand for 3 minutes with wash solution in the wells. Remove wash solution from wells and repeat two additional times.

Other wash solutions may be used, however, the solution must not contain chelators of divalent metal cations, i.e., no EDTA.

 Add 100 µL/well of KPL BluePhos[®] Substrate for detection. As the color develops, tap gently to mix. Incubation times will vary depending on the assay.

> Other substrates are available for AP detection in ELISA. SeraCare also offers KPL pNPP for colorimetric ELISA detection. For chemiluminescent detection, KPL PhosphaGLOTM or KPL PhosphaGLOTM Reserve are available.

> See RELATED PRODUCTS on page 22 for more information on AP substrates for ELISA.

- 4. Stop reaction by adding 100 µl of KPL BluePhos Stop Solution (See RELATED PRODUCTS) or 2.5% EDTA to each well.
- Read plate using a plate reader at the appropriate wavelength (630 nm for KPL BluePhos) or a luminometer (chemiluminescent detection).

TROUBLESHOOTING GUIDE KPL HisDetector[™] Western Blot Protocols

Use the dot blot protocol on page 13 to troubleshoot Western Blot protocols. The dot blot protocol saves time by omitting electrophoresis and transfer steps.

Problem 1: No Signal

Pos	ssible Cause	Corrective Measure
•	The membrane was blocked in an incompatible block solution, such as those containing milk- based or phosphorylated proteins.	Block membrane in 1% BSA/TBST for KPL Nickel-HRP or KPL 1X Detector Block Solution for KPL Nickel-AP.
•	Inhibition of nickel-NTA binding to His-tagged protein.	Ensure buffers or reagents do not contain chelators such as EDTA, imidazole, or other divalent metal cations.
•	Presence of an inhibitor of enzymatic activity.	Ensure buffers do not contain inhibitors such as sodium azide or phosphatase inhibitors. Sodium azide is a potent inhibitor of HRP activity.
•	Insufficient sample loaded in	Determine total protein concentration.
	lane or low level of expression of His-tagged protein.	Using protein molecular markers, estimate level of protein expression from a Coomassie stain.
•	Lack of His-tag fusion with recombinant protein.	Ensure the recombinant protein is His- tagged.
•	Incomplete transfer of proteins to membrane.	Use a pre-stained protein ladder to establish level of transfer.
•	Incorrect orientation of membrane during exposure to film.	Check that the correct orientation of the membrane was maintained during the assay, that it was not placed "wrong" side down on film.
•	Reagent omitted or improperly prepared.	Check that all procedures were carried out correctly.

Problem 2: Weak Signal

Possible Cause	Corrective Measure
• The membrane was blocked in an incompatible block solution, such as those containing milk- based or phosphorylated proteins.	Block membrane in 1% BSA/TBST for KPL Nickel-HRP or KPL 1X Detector Block Solution for KPL Nickel-AP.

•	Insufficient sample loaded or low expression of His-tagged protein.	Determine total protein concentration. Using protein molecular markers, estimate level of protein expression from a Coomassie stain.
•	Short exposure of membrane to film	Expose the film to the membrane for extended time, such as 10 minutes.
•	His-tag region of protein is not exposed.	Add ß-mercaptoethanol to denature protein.
		Heat sample prior to loading on gel.
		After transfer, use chaotropic agents (6M urea or 3M potassium thiocyanate ¹) to increase accessibility of the nickel binding region.
•	Assay system may require	Titrate nickel conjugate concentration.
	greater quantities of KPL Nickel-HRP or AP conjugate or longer exposure to conjugate.	Increase the incubation time of nickel conjugate.
		Increase the time of film exposure.
•	Incomplete transfer of proteins to membrane.	Use a pre-stained protein ladder to establish level of transfer.

Problem 3: Excessive Signal

	_	
Po	ssible Cause	Corrective Measure
 Bands "blasting" their signal of bands not visible due to excessive signal. 	Bands "blasting" their signal or bands not visible due to	Titrate total amount of protein loaded on the gel.
	excessive signal.	Reduce concentration of Nickel-NTA conjugate.
		Reduce substrate incubation time or film exposure.
•	Film exposure time is too long.	Decrease film exposure time.

Problem 4: Non-Specific Binding of Nickel Conjugate

Possible Cause		Corrective Measure
•	Lane is overloaded with protein.	Load less amount of protein sample. Decrease film exposure time.
•	KPL Nickel-HRP or AP is too concentrated.	Optimize KPL Nickel-HRP or AP concentration using serial dilutions.

Problem 5: High Background

Possible Cause		Corrective Measure
•	Insufficient washing.	Increase wash times if necessary.
		Ensure blot is covered with 1X TBST.
•	Insufficient blocking.	Use blocking solutions recommended in appropriate western blot protocol.
		Ensure blot is covered with blocking solution.
•	Endogenous phosphatase or peroxidase activity in blocking solution.	Use blocking solutions recommended in appropriate western blot protocol.
•	Excess substrate on the membrane during film exposure.	Remove excess substrate from membrane before exposure to the film by touching membrane to a piece of filter paper.

Problem 6: Detection of Multiple Protein Bands Containing His-tags

Possible Cause		Corrective Measure
•	Insufficient amount of reducing agent.	Titrate concentration of reducing agent.
•	Degradation of protein of interest due to protease activity.	Consider use of protease inhibitors at the lysis step.

TROUBLESHOOTING GUIDE KPL HisDetector[™] ELISA Protocols

Problem 1: No Signal

Pos	sible Cause	Corrective Measure	
•	The ELISA plate was blocked in an incompatible block solution, such as those containing milk-based or	Block the ELISA plate in 1% BSA/TBST for KPL Nickel-HRP or KPL 1X Detector Block Solution for KPL Nickel-AP.	
•	phosphorylated proteins. Inhibition of nickel-NTA binding to His-tagged protein.	Ensure buffers or reagents do not contain EDTA, imidazole, or other metal chelators.	
•	Presence of an inhibitor of HRP or AP in assay.	Use buffers without chelators such as EDTA, imidazole, or other divalent metal cations.	
•	Reagent omitted or improperly prepared.	Check that all procedures were carried out correctly.	
•	Insufficient amount of sample plated in each well	Optimize the protein concentration needed via several titrations.	
•	Lack of His-tag fusion on the recombinant protein.	Ensure the recombinant protein contains a His-tagged portion.	
Problem 2: Weak Signal			
Pos	sible Cause	Corrective Measure	
•	Insufficient quantity of target protein plated or low level of expression of His-tagged protein.	Increase amount of protein coated on plate.	
•	His-tag region of protein is not well exposed.	Add ß-mercaptoethanol or DTT to denature protein. Heat sample prior to coating on plate.	
•	Assay system may require higher levels of Nickel conjugate.	Titrate nickel conjugate concentration. Increase nickel conjugate incubation time.	
Pro	blem 3: High Background	1	
Pos	sible Cause	Corrective Measure	
•	Excess protein coated on ELISA plate.	Reduce concentration of His-tagged protein coated on plate.	
•	KPL Nickel-HRP or AP conjugate is too concentrated.	Optimize KPL Nickel-HRP or AP concentration using serial dilutions.	
	conjugate is too concentrated.	g	
•	Insufficient washing.	Increase soak time for each wash step or add additional wash steps.	

REFERENCES

Jin, L. et al. (1995) *Analytical Biochem.* 229, 54-60. Peterson, D. (1996) *Australian Biotechnology.* 6, 103-106. Jin, L. & Peterson, D. (1995) *Arch. Biochem. Biophys.* 323, 47-53.

RELATED PRODUCTS

Western Blotting Products	<u>Cat. No.</u>	<u>Size</u>
KPL Detector TM Block (5X)	5920-0004 (71-83-00)	240 mL
KPL TMB Membrane HRP Substrate	5420-0029 (50-77-18)	100 mL
KPL LumiGLO HRP Chemiluminescent Substrate	5430-0040 (54-61-00)	240 mL
KPL LumiGLO Reserve HRP Chemiluminescent Substrate	5430-0049 (54-71-00)	2400 cm ²
KPL PhosphaGLO AP Substrate	5430-0055 (55-60-04)	100 mL
KPL PhosphaGLO Reserve AP Substrate	5430-0053 (55-60-02)	100 mL
KPL BCIP/NBT AP Substrate	5420-0038 (50-81-18)	100 mL

ELISA Products	<u>Cat. No.</u>	<u>Size</u>
KPL 10X Coating Solution	5150-0014 (50-84-00)	50 mL
KPL ABTS ELISA HRP Substrate	5120-0046 (50-66-18)	100 mL
KPL ABTS ELISA HRP Substrate	5120-0032 (50-62-00)	600 mL
KPL ABTS HRP Stop Solution	5150-0017 (50-85-01)	200 mL
KPL SureBlue TMB ELISA HRP Substrate	5120-0075 (52-00-01)	100 mL
KPL SureBlue Reserve TMB ELISA	5120-0081 (53-00-01)	100 mL
KPL TMB Stop Solution	5150-0020 (50-85-05)	400 mL
KPL pNPP ELISA AP Substrate System	5120-0056 (50-80-00)	500 mL
KPL pNPP ELISA AP Substrate (100 tablets)	5120-0057 (50-80-01)	5 mg tabs
KPL BluePhos ELISA AP Substrate	5120-0059 (50-88-00)	600 mL
KPL BluePhos Stop Solution (10X)	5150-0026 (50-89-00)	200 mL

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HisDetectorTM nickel conjugates are produced with components protected by U.S. Patents 6,800,728, 5,679,778, 5,420,285, 5,753,520 and 5,206,370.

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SeraCare Life Sciences

508.244.6400 • 800.676.1881 Toll Free • 508.634.3334 Fax www.seracare.com

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