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A Geno Technology, Inc. (USA) brand name

# *femto*LUCENT<sup>™</sup> PLUS-HRP

## Chemiluminescence Detection System for Horseradish Peroxidase

(Cat. # 786-003, 786-056, 786-081, 786-10, 786-10T)



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## **INTRODUCTION**

femtoLUCENT<sup>T</sup> PLUS-HRP kit is based on our ultra sensitive Luminol substrate that produces chemiluminescence upon reaction with horseradish peroxidase. The chemiluminescence light emission can be recorded by a short exposure to autoradiography films.

femtoLUCENT<sup>™</sup> PLUS-HRP reagents are available in three sizes suitable for 25, 50 and 125 blots as each 4ml of working solution is suitable for 1 mini blot (8 x 7.5cm)

In addition, femtoLUCENT<sup>TM</sup> PLUS-HRP is also supplied in a kit format, containing our nonanimal protein blocking agent (NAP-BLOCKER<sup>TM</sup>) and wash buffer (femto-TBST<sup>TM</sup>). The femtoLUCENT<sup>TM</sup> PLUS-HRP kits allow detection of low femtogram levels ( $10^{-15}$ ) of antigens with low noise (signal/background) ratio. The kit reagents are sufficient for 25 mini blots or 1,500cm<sup>2</sup> of PVDF or nitrocellulose membrane. The trial size (Cat. # 786-10T) is suitable for 5 mini blots or 300cm<sup>2</sup>.

| <b>ITEM(S)</b> | <b>SUPPLIED</b> |
|----------------|-----------------|
|----------------|-----------------|

| Cat. #  | Size* | NAP-BLOCKER <sup>™</sup><br>[2X] | femto<br>TBST<br>[10X] |       | Peroxide<br>Solution |
|---------|-------|----------------------------------|------------------------|-------|----------------------|
| 786-056 | 50    | -                                | -                      | 100ml | 100ml                |
| 786-081 | 125   | -                                | -                      | 250ml | 250ml                |
| 786-003 | 25    | -                                | -                      | 50ml  | 50ml                 |
| 786-10  |       | 250ml                            | 250ml                  | 50111 | 30111                |
| 786-10T | 5     | 125ml                            | 50ml                   | 10ml  | 10ml                 |

\* For mini blots with average size of 8 x 7.5cm or 60cm<sup>2</sup>.

## **STORAGE CONDITIONS**

Shipped at ambient temperature. Upon arrival, store the kit components at 4°C, protected from light. When stored and used properly, the kit is stable for 1 year.

## **ITEMS NOT SUPPLIED**

- Primary antibody
- Secondary antibodies, HRP-conjugates.

#### **PREPARATION BEFORE USE**

- Preparation of Working Detection Solution: Allow the solutions to warm to room temperature before use. For each mini blot membrane (~60cm<sup>2</sup>), mix 2.0ml of femtoLUCENT<sup>™</sup> Luminol Solution-A and 2.0ml femtoLUCENT<sup>™</sup> Peroxide Solution-B.
- Preparation of 1X femto-TBST: Dilute the appropriate volume of supplied 10X femto-TBST to 1X with DI Water (e.g. Take 10ml of 10X femto-TBST and add 90ml DI Water to make it 1X).
- 3. Preparation of 1X NAP-BLOCKER<sup>®</sup>: Use aseptic techniques for handling NAP-BLOCKER<sup>®</sup>. Allow the supplied 2X NAP-BLOCKER<sup>®</sup> bottle to come to room temperature and then gently shake to mix. Dilute the appropriate volume of 2X NAP-BLOCKER<sup>®</sup> 1:1 with 1X femto-TBST (e.g. Take 10ml of 2X NAP-BLOCKER<sup>®</sup> and add 10ml of 1X femto-TBST).

#### **PROTOCOL FOR REAGENT ONLY**

- 1. Prepare the Working Detection Solution as described above.
- 2. Block and probe your membrane with appropriate blocking agents and antibodies as per your regular protocols.
- 3. Following the final wash steps, add 4ml Working Detection Solution for a mini blot and incubate for 3-5 minute at room temperature with gentle shaking.
- Drain the detection reagent and wrap the membrane in saran wrap or clear plastic wrap and expose to an autoradiography film.
  NOTE: Do NOT wash or rinse the membrane after addition/removal of the working detection solution.

#### **PROTOCOL FOR FEMTOLUCENT<sup>™</sup> PLUS-HRP KIT**

- Blocking: After the electrophoretic transfer of the protein to an appropriate membrane (e.g. PVDF or Nitrocellulose), block the membrane by immersing in 10ml 1X NAP-BLOCKER<sup>™</sup>. Incubate the blot (membrane) in the blocking buffer for a minimum of 60 minutes at room temperature with gentle shaking on an orbital shaker.
- Primary Antibody Treatment: Dilute the primary antibody in an appropriate volume (≤5ml) of 1X NAP-BLOCKER<sup>™</sup>. Incubate the membrane in the diluted primary antibody for 1-2 hours at room temperature, with gentle shaking.

**NOTE:** Determine the optimal dilution of the primary antibody in separate experiments or follow the manufacturer's instructions.

- Washing: Rinse the membrane with ~10ml 1X *femto*-TBST then wash three times with ~10ml 1X *femto*-TBST buffer for 10 minutes each at room temperature with gentle shaking.
- 4. Secondary Antibody Treatment: Dilute the HRP-conjugated secondary antibody in an appropriate volume (≤5ml) of 1X NAP-BLOCKER<sup>™</sup> at a 1:5,000 to 1:100,000 dilution. Incubate the membrane in the diluted secondary antibody for 1-2 hours at room temperature with gentle shaking.

**NOTE:** Determine the optimal dilution of the secondary antibody in separate experiments.

- Washing: Rinse the membrane with ~10ml 1X *femto*-TBST then wash three times with ~10ml 1X *femto*-TBST buffer for 10 minutes each at room temperature with gentle shaking.
- 6. **Chemiluminescence Reaction**: Incubate the membrane in the 4ml Working Detection Solution for 3-5 minute at room temperature with gentle shaking.
- 7. Drain the detection reagent and wrap the membrane in saran wrap or clear plastic wrap and expose to an autoradiography film.

**NOTE:** Do NOT wash or rinse the membrane after addition/removal of the working detection solution.

## **REDEVELOPING THE MEMBRANE**

The membrane can be redeveloped within a day or two, provided that the detection reagents are removed from the membrane within 30-60 minutes of the first developing procedure. After each developing procedure, wash the membrane with 50ml TBS with Tween-20. Keep the membrane moist and at 4°C. Redevelop the membrane according to the protocol above and expose the autoradiography film.

## TROUBLESHOOTING

## Uniform High Background

| Suggested Cause   | Resolution/ Precaution   |
|---|--|
| Concentration of antibody<br>too high                                 | Reduce the concentration of primary and/ or secondary antibodies as high concentration can result in a high background.  |
| Interference from<br>incompatible blocking agent                      | Investigate a different blocking agent, such as non animal protein blocking agents.  |
| Antibodies cross-react with<br>proteins present in blocking<br>agents | Investigate a different blocking agent, such as non<br>animal protein blocking agents.<br>Avoid milk based blocking agents when probing with<br>avidin/biotin systems. Milk contains biotin.   |
| Non-specific sites<br>insufficiently blocked                          | Optimize the blocking buffer and conditions, including<br>amount and type of blocking protein (agent) and length<br>and temperature of blocking step.<br>Add Tween <sup>®</sup> 20 to the blocking agent, if detergent is<br>not already present. Final concentration of 0.05%.<br>Incubate with antibodies in blocking agent containing<br>0.05% Tween <sup>®</sup> 20. |
| Washing steps insufficient  | Increase volume and length of wash steps.<br>Use wash buffers with Tween <sup>®</sup> 20, such as out femto-<br>TBST <sup>™</sup> or femto-TBST <sup>™</sup> Wash Buffers.   |
| Membrane exposed too long to film                                     | Reduce the exposure time.  |
| Membrane issues   | Membranes not wetted correctly, check<br>manufacturer's instructions.<br>Membrane inadvertently dried out during procedure.<br>Use orbital shaking or rocking with all incubation steps.<br>Handle membrane carefully, do not touch with<br>exposed skin.  |
| Bacterial or other contamination                                      | Prepare fresh buffers.   |

Blotchy or Speckled High Background

| Suggested Cause   | Resolution/ Precaution  |
|---|---|
| Concentration of antibody<br>too high                                 | Reduce the concentration of primary and/ or secondary antibodies as high concentration can result in a high background.   |
| Interference from<br>incompatible blocking<br>agent                   | Investigate a different blocking agent, such as non animal protein blocking agents.   |
| HRP conjugate may have aggregated                                     | Filter through a 0.2µm filter.<br>Use new conjugate.  |
| Antibodies cross-react with<br>proteins present in<br>blocking agents | Investigate a different blocking agent, such as non animal<br>protein blocking agents.<br>Avoid milk based blocking agents when probing with<br>avidin/biotin systems. Milk contains biotin.  |
| Non-specific sites<br>insufficiently blocked                          | Optimize the blocking buffer and conditions, including<br>amount and type of blocking protein (agent) and length<br>and temperature of blocking step.<br>Add Tween <sup>®</sup> 20 to the blocking agent, if detergent is not<br>already present. Final concentration of 0.05%.Incubate<br>with antibodies in blocking agent containing 0.05%<br>Tween <sup>®</sup> 20. |
| Washing steps insufficient  | Increase volume and length of wash steps.<br>Use wash buffers with Tween <sup>®</sup> 20, such as out femto-<br>TBST <sup>™</sup> or femto-TBST <sup>™</sup> Wash Buffers.  |
| Membrane exposed too<br>long to film                                  | Reduce the exposure time.   |
| Membrane issues   | Membranes not wetted correctly, check manufacturer's<br>instructions.<br>Membrane inadvertently dried out during procedure.<br>Use orbital shaking or rocking with all incubation steps.<br>Handle membrane carefully, do not touch with exposed<br>skin.   |
| Bacterial or other contamination                                      | Prepare fresh buffers.  |

| Suggested Cause | Resolution/ Precaution   |  |
|-----------------|--|--|
| Dirty equipment | Ensure all equipment is free of contaminants.<br>Ensure no residual gel pieces are present on the<br>membrane. |  |

## Weak or No Signal

| Suggested Cause                           | Resolution/ Precaution   |
|---|--|
| Improper transfer of proteins to membrane | Ensure correct protein transfer by staining the membrane<br>with a suitable, reversible membrane stain. We recommend<br>Swift <sup>™</sup> Membrane Stain (Cat. No. 786-677).<br>For poor transfer of high molecular weight proteins, use our<br>High Molecular Weight Transfer Buffer (Cat. No. 786-423).<br>Ensure the transfer sandwich and apparatus is assembled<br>correctly and electrodes are correctly orientated.<br>Avoid over heating during transfer. |
| Poor binding of protein to membrane       | Add 20% methanol to transfer buffer for improved binding.<br>For low molecular weight proteins, reduce transfer time or<br>use a membrane with a smaller pore size to prevent proteins<br>passing through.   |
| Concentration of antibody too high        | Reduce the concentration of primary and/ or secondary<br>antibodies as high concentration can result in the signal<br>generating too quickly and fading away before development.   |
| Concentration of antibody too low         | Increase antibody concentration to overcome possible poor affinities.  |
| Antigen levels too low                    | Load more proteins on the initial gel.   |
| Blocking agent binds<br>antigen           | Optimize blocking agent type and concentration.  |
| HRP activity inhibited by sodium azide    | Avoid using buffers that use sodium azide as a preservative.   |
| Exposure time too<br>short                | Extend the film exposure time.   |

| Suggested Cause              | Resolution/ Precaution   |  |
|------------------------------|--|--|
| Detection substrate inactive | Ensure the substrate(s) shelf life has not expired.<br>Cross reaction between the 2/3 component systems may<br>have occurred.<br>Check for activity by preparing substrate and, in a dark roo<br>add a small amount of conjugate. If active a blue light shou<br>appear. |  |
| Excessive stripping          | If the membrane has been stripped, antigen sites may have<br>been destroyed.<br>Use mild stripping conditions, we recommend Western<br>ReProbe <sup>™</sup> (Cat. No. 786-119).<br>Limited the number of times a membrane is reprobed.                                   |  |

## Non-Specific Bands

| Suggested Cause                    | Resolution/ Precaution   |  |
|------------------------------------|--|--|
| Concentration of antibody too high | Reduce the concentration of primary and/ or secondary antibodies as high concentration can result in non-specific binding. |  |
| Presence of SDS                    | Thoroughly wash blot after transfer.<br>Do not use SDS in development steps.   |  |

## Diffuse Bands

| Suggested Cause                    | Resolution/ Precaution   |  |
|------------------------------------|--|--|
| Concentration of antibody too high | Reduce the concentration of primary and/ or secondary antibodies as high concentration can result in non-specific binding. |  |
| Protein concentration too high     | Reduce the amount of protein loaded  |  |

## Ghost/ Hollow Bands or Brown/Yellow Bands on Membrane

| Suggested Cause                    | Resolution/ Precaution   |
|------------------------------------|--|
| Concentration of antibody too high | Reduce the concentration of primary and/ or secondary antibodies as high concentration can result in rapid consumption of the substrate. |

#### Blank Areas

| Suggested Cause | Resolution/ Precaution  |
|-----------------|---|
| Incomplete      | Ensure correct protein transfer by staining the membrane with a suitable, reversible membrane stain. We recommend Swift <sup>™</sup> Membrane Stain (Cat. No. 786-677). |
| transfer of     | Ensure the transfer sandwich and apparatus is assembled correctly and electrodes are correctly orientated.  |
| proteins        | Avoid over heating during transfer.   |

#### STRIPPING AND RE-PROBING MEMBRANE

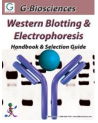
The developed membrane can be stripped and re-probed with any other antibody by using a suitable stripping buffer. G-Biosciences Western-Re-Probe<sup>™</sup> Buffer (5X) is recommended for stripping and re-probing procedures.

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