About us

Cyanagen is a biotech company located in Bologna, dedicated to research, development and production of reagents for molecular diagnostic since 2003 and one of the leading companies in the field of reagents for Western blotting and Elisa. The main product lines are focused on chemiluminescence and fluorescent dyes for biological analysis, genomics, proteomics and chemical sensors. They are based on Cyanagen internationally patented technologies and achieve outstanding performance in terms of sensitivity and stability. The products are extremely versatile and perfectly suited to the latest analytical instrumentation. These products are also available as OEM.

Cyanagen s.r.l. has a certified Quality System

ISO 9001-2008 QUALITY CERTIFIED

ECL substrate for Western Blotting

WESTAR IS INTENDED FOR RESEARCH USE ONLY AND SHALL NOT BE USED IN ANY CLINICAL PROCEDURES OR FOR DIAGNOSTIC PURPOSES.

www.cyanagen.com
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1. Introduction

The peroxidase-catalyzed oxidation of luminol produces a weak flash of light at 425 nm. The incorporation of an electron transfer mediator into the buffer forces the flash signal into a glow and greatly improves the analytical characteristics of the reaction in terms of increased signal intensity and duration.\textsuperscript{1,2} Recent works\textsuperscript{3+6} have shown that, by addition of a suitable acylation catalyst, a further large increase in light output is observed.

WESTAR detection reagents are non-isotopic, luminol-based chemiluminescence substrate, designed for the chemiluminescent detection of immobilized proteins and immobilized nucleic acids conjugated with horseradish peroxidase (HRP). WESTAR is intended for research use only, and shall not be used in any clinical procedures, or for diagnostic purposes.

References:

Storage/expiry
One year at room temperature (25° C).

WESTAR product line

<table>
<thead>
<tr>
<th>WESTAR</th>
<th>SUN</th>
<th>NOVA 2.0</th>
<th>ETA C 2.0</th>
<th>ETA C ULTRA 2.0</th>
<th>SUPERNOVA</th>
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</thead>
<tbody>
<tr>
<td>Signal intensity</td>
<td>Standard</td>
<td>Medium</td>
<td>High</td>
<td>Very High</td>
<td>Extreme</td>
</tr>
<tr>
<td>Signal duration</td>
<td>Very long</td>
<td>Long</td>
<td>Good</td>
<td>Moderate</td>
<td>Short</td>
</tr>
<tr>
<td>Protein quantity</td>
<td>High abundance</td>
<td>High abundance</td>
<td>Medium abundance</td>
<td>Low abundance</td>
<td>Very-low abundance</td>
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</table>
2. Components and other materials required

**Kit components**

- **Solution A**: Luminol/enhancer solution (amber bottle)
- **Solution B**: Peroxide solution (white bottle)

**Other required solutions**

<table>
<thead>
<tr>
<th>Solution</th>
<th>Preparation</th>
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<tr>
<td><strong>Running Buffer</strong></td>
<td><strong>For 1L of 10x Running Buffer (stock):</strong></td>
</tr>
<tr>
<td></td>
<td>- 30.3 g TRIS (250mM)</td>
</tr>
<tr>
<td></td>
<td>- 144.0 g Glycine (1.9M)</td>
</tr>
<tr>
<td></td>
<td>- 10.0 g SDS (1% w/v)</td>
</tr>
<tr>
<td></td>
<td>- Dilute to 1L with distilled water</td>
</tr>
<tr>
<td></td>
<td><strong>For 1L of Running Buffer:</strong></td>
</tr>
<tr>
<td></td>
<td>- 100mL of <strong>10x Transfer Buffer</strong></td>
</tr>
<tr>
<td></td>
<td>- Dilute to 1L with distilled water</td>
</tr>
<tr>
<td><strong>Transfer Buffer</strong></td>
<td><strong>For 1L of 10x Transfer Buffer (stock):</strong></td>
</tr>
<tr>
<td></td>
<td>- 30.3 g TRIS (250mM)</td>
</tr>
<tr>
<td></td>
<td>- 144.0 g Glycine (1.9M)</td>
</tr>
<tr>
<td></td>
<td>- Dilute to 1L with distilled water</td>
</tr>
<tr>
<td></td>
<td><strong>For 1L of Transfer Buffer:</strong></td>
</tr>
<tr>
<td></td>
<td>- 100mL of <strong>10x Transfer Buffer</strong></td>
</tr>
<tr>
<td></td>
<td>- 200mL of methanol</td>
</tr>
<tr>
<td></td>
<td>- Dilute to 1L with distilled water</td>
</tr>
<tr>
<td><strong>TBS-T Buffer</strong></td>
<td><strong>For 1L of 10x TBS Buffer (stock):</strong></td>
</tr>
<tr>
<td></td>
<td>- 24.23 g TRIS-HCl (20mM)</td>
</tr>
<tr>
<td></td>
<td>- 80.06 g NaCl (136mM)</td>
</tr>
<tr>
<td></td>
<td>- Dilute to 800mL with distilled water</td>
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<tr>
<td></td>
<td>- Add NaOH 1M until pH is about 7.6</td>
</tr>
<tr>
<td></td>
<td>- Dilute to 1L with distilled water</td>
</tr>
<tr>
<td></td>
<td><strong>For 1L of TBS-T Buffer:</strong></td>
</tr>
<tr>
<td></td>
<td>- 100 ml of <strong>10x TBS Buffer</strong></td>
</tr>
<tr>
<td></td>
<td>- While stirring add 1 mL Tween-20</td>
</tr>
<tr>
<td></td>
<td>- Dilute to 1L with distilled water</td>
</tr>
<tr>
<td><strong>Blocking Buffer</strong></td>
<td><strong>With 5% non-fat dried milk:</strong></td>
</tr>
<tr>
<td></td>
<td>- 5 g Non-fat dried milk</td>
</tr>
<tr>
<td></td>
<td>- Dissolve in 100 ml <strong>1x TBS-T Buffer</strong></td>
</tr>
<tr>
<td></td>
<td><strong>With 5% BSA:</strong></td>
</tr>
<tr>
<td></td>
<td>- 5 g BSA (Cohn fraction V)</td>
</tr>
<tr>
<td></td>
<td>- Dissolve in 100 ml <strong>1x TBS-T Buffer</strong></td>
</tr>
<tr>
<td><strong>Ponceau staining solution</strong></td>
<td><strong>For 100mL of 10x Ponceau staining solution (stock):</strong></td>
</tr>
<tr>
<td></td>
<td>- Dissolve 0.5 g Ponceau S in 1.0 ml glacial acetic acid</td>
</tr>
<tr>
<td></td>
<td>- Dilute to 100 ml with distilled water</td>
</tr>
<tr>
<td></td>
<td>- Wrap bottle with foil to protect solution from light</td>
</tr>
<tr>
<td></td>
<td><strong>For 1L of Ponceau staining solution:</strong></td>
</tr>
<tr>
<td></td>
<td>- 100 ml of <strong>10x Ponceau S staining solution</strong></td>
</tr>
<tr>
<td></td>
<td>- Dilute to 100 ml with distilled water</td>
</tr>
</tbody>
</table>

**Compatible imaging devices**

- Westar R (HiTech Cyanagen)
- ImageQuant™LAS-4000/Mini (GE Healthcare)
- DIAS-II (SERVA)
- ChemiDocXRS and VersaDoc (BIO-RAD)
- ChemiImager (Alpha Innotech)
- Image Station 2000/40000MM (Kodak)
- FOTO/Analyst Luminary/FX Systems (Fotodyne)
- LAS-3000 (Fujifilm)
- UVichemi and UVIprochemi (UVItec Ltd.)
- G:BOX/GeneGnome (Syngene)
- Odyssey FC (LI-COR)
3. Perform SDS-PAGE

I. Prepare fresh **Running Buffer**.
II. Load the gels being sure to keep a tight seal between the gel-cast and the gasket.
III. Pour the running buffer into the middle of the gels and check for leaks.
IV. Pour the rest of the running buffer into the bottom of the running tank.
V. Remove combs and use a pipette to clean away any unpolymerized acrylamide.
VI. Load a proper **prestained MW standard** in one lane.
VII. Load samples into the rest of the wells and fill any empty well with sample buffer.
VIII. Run at 90÷130 V constant voltage until the dye front reaches the bottom of the gel. If the current is too high band smiling and smearing (diffuse band) are commonly seen effects.

4. Prepare transfer membrane

If using **nitrocellulose membrane** place into distilled water slowly, with one edge at a 45° angle. If inserted too quickly into the water, air gets trapped and protein will not transfer onto these areas. Once wet, equilibrate the membrane in **Transfer Buffer** for 15 min.

If using **PVDF membrane** activate it with methanol for 30 seconds. Rinse with distilled water and equilibrate in **Transfer Buffer** for 15 min.

- For proteins >15 kDa use membrane pore size 0.45 mm
- For proteins <15 kDa use membrane pore size 0.2 mm

**NOTE:** Low molecular weight proteins (< 15kDa) are sometimes transferred through nitrocellulose membranes, therefore may be not visible on the blot. PVDF membrane has higher protein binding capacity than nitrocellulose membrane and is recommended for best detection sensitivity.
5. Transfer to membrane

I. Wet four filter papers in **Transfer Buffer**.

II. Assemble the transfer sandwich in a tray large enough to hold the plastic transfer cassette. Fill with **Transfer Buffer** so that the cassette is covered.

III. Place the first foam pad onto the **black side** of the transfer cassette then place two pre-wetted filter papers on the top of it.

IV. Place the gel and moisten its surface with **Transfer Buffer**.

V. Place pre-wetted membrane directly on the top side of the gel, then gently remove all air bubbles. The proteins will transfer as soon as the gel is placed on the membrane, its repositioning can generate a smeared image.

VI. Place another two pre-wetted filter paper over the membrane and remove all air bubbles.

VII. Complete the assembly by placing the last foam pad and locking the top half of the transfer cassette.

VIII. Fill the transfer tank with **Transfer Buffer** and place the transfer cassette.

IX. Put a frozen cooling unit into the transfer tank and surround it with ice in a polystyrene box.

X. Run the transfer with the following settings:
   - **Wet transfer**: 80÷100 V for 30÷60min.
   - **Semi-dry transfer**: 15÷25 V for 20÷30min.

XI. When transfer is complete, remove the membrane and mark its orientation by cutting a corner.

XII. Wash the membrane twice with distilled water.

**Constant voltage or current during transfer?**

The buffer composition changes as salts are eluted from the gels, resulting in an increase in current and a drop in resistance. A transfer using constant current leads to decrease in voltage as well as resistance (I=V/R). Therefore, the use of constant voltage provides the best driving force during transfer. However, when current reaches over 500mA in constant voltage setting cooling the gel is crucial for preventing joule heating in the tank.
6. Membrane staining (optional)

I. Stain the membrane with protein side up using Ponceau staining solution for 5 minutes at RT to check transfer efficiency.
II. Rinse the membrane in distilled water until protein bands are distinct.
III. Scan the membrane if desired.
IV. Completely destain the membrane by immersing it for 10 min in a large volume of distilled water.
V. Re-activate PVDF membrane with methanol then wash in TBS-T Buffer.

NOTE:
- The background staining tends to be high with some dyes while Ponceau staining solution gives a very clean pattern.
- Re-activate PVDF membrane after staining.
- The LOD for Ponceau staining solution is 250 ng of protein.

7. Blocking the membrane

I. Place the membrane with protein side up into a fresh tray with your choice of Blocking Buffer.
II. Incubate the membrane in Blocking Buffer for 30÷60 minutes with gentle agitation on a rocker/shaker. A maximum blocking time of 2 hours at RT should not be exceeded. Blocking for too long can result in antigen masking and loss of protein.
III. Rinse the membrane twice with TBS-T Buffer.

NOTE: Add 3% non-fat dry milk in TBS-T Buffer when dilute Ab to reduce non specific bindings. Milk contains many proteins which bind to the membrane. So, after transfer, proteins contained in the milk bind to the membrane and fill a lot of potential non specific sites. After this, when you incubate with your antibody, it binds to the antigen and has less possibilities of non specific binding. If you are working with anti-phosphoproteins or with biotinylated antibodies the adding of milk is not appropriate. Use 5% BSA instead.
8. Antibody incubation

I. Dilute the primary antibody in fresh TBS-T Buffer to the suggested primary antibody dilution (see table below).

II. Incubate the membrane with protein side up in the primary antibody solution for 1 to 2 hours at RT. To increase sensitivity, try an overnight incubation at 4°C with agitation on a rocker. Make sure the membrane is completely covered with TBS-T Buffer with primary Ab.

III. Wash the membrane with protein side up 4 times for 3 to 5 minutes each with TBS-T Buffer with gentle agitation on a shaker. After each washing place the membrane on a new clean tray with fresh TBS-T buffer.

IV. Dilute the secondary Ab in fresh TBS-T Buffer to the suggested secondary antibody dilution (see table below).

V. Incubate the membrane with protein side up for 30 minutes to 1 hour at RT. Increasing the incubation time of the secondary antibody usually leads to higher background.

VI. Wash the membrane with protein side up 4 times for 3 to 5 minutes each with TBS-T Buffer with gentle agitation on a shaker. After each washing place the membrane on a new clean tray with fresh TBS-T buffer.

IMPORTANT: Optimal Ab dilutions may vary between different applications and depend on quality and affinity for the target protein. It is crucial to optimize both primary and secondary Ab dilutions for best results with high signal and low background. Optimal Ab dilutions can be determined by Dot-Blot assay.

Suggested Ab dilutions for WESTAR NOVA 2.0

- Primary Ab ................ from 1:500 to 1:5,000
- Secondary Ab ............... from 1:20,000 to 1:100,000
9. Chemiluminescent detection

I. For reproducible performance allow the detection solutions to equilibrate to RT before using.  
II. Prepare Westar working solution (Westar WS) by mixing properly each reagent in a 1:1 ratio. For best results prepare WS immediately before use. Do not contaminate the solutions with the same pipette tips.  
III. Remove the membrane from its tray of TBS-T Buffer and rinse the membrane twice with TBS-T Buffer and keep it in TBS until the incubation with Westar WS.  
IV. Use 0.1 ml of Westar WS per cm² of membrane.  
V. Allow the excess buffer to run off from a corner. Do not let the membrane dry out. Just pipette the volume required directly onto the membrane with protein side up and incubate for 1.5 min ensuring that the entire surface is covered.  
VI. Acquire the signal with autoradiography film or imaging devices. For an unknown signal, try to expose 15 s, 30 s, 1 min and 5 min to start with.

Autoradiography film vs. imaging devices

Nowadays, Western Blotting is used either for absolute quantification (in combination with a calibration curve of the recombinant protein of known concentration) or for quantification of samples relative to a control sample. Through the development of new technologies most imagers offer a wide dynamic range (3÷5 orders of magnitude) generating a high-quality image compared with the limited linear dynamic range of film (1.5 orders of magnitude). This means that is possible to quantify both strong and weak signals on the same blot with reliable results. Instead, on film strong signals get saturated resulting in a wrong quantitation.
10. Troubleshooting

High membrane background

High concentration of Ab. Further dilute primary and secondary Ab. Follow suggested Ab dilutions.

Inefficient blocking. Increase Tween-20 in TBS-T Buffer (0.1%÷0.5% v/v). Use 5% non-fat dried milk as blocking buffer if possible.

Insufficient washing. Increase both the volume, length and number of wash steps. Always use sufficient volumes to submerge the membrane.

Primary antibody is not specific for the protein of interest. Use monospecific or antigen affinity purified antibodies. Always incubate your primary antibody at 4°C overnight and not at room temperature. Reduce NaCl in TBS-T Buffer (100mM÷350mM). Use monospecific or antigen affinity purified Ab.

Non-specific binding of secondary antibody. Confirm the secondary is specific by omitting the primary and running a secondary only blot. If bands develop choose an alternative secondary antibody.

Incompatible blocking agent. Non-fat dry milk contains endogenous biotin and is incompatible with avidin/streptavidin systems. Substitute with 5% BSA.

Poor quality of antibodies. Quality and age of primary and secondary antibody may lead to background problems.

Poor handling of membrane. Be sure to handle the membrane only with clean plastic tweezers and non-powdered gloves.

Contaminated buffer solutions. Check buffers for particulate or bacterial contaminate. Replace old buffers.

Irregular black spots

Air bubble trapped in membrane. Remove air bubbles by gently rolling a clean pipette or a test-tube during sandwich assembling.

Unevenly hydrated membrane. Make sure that the membrane is fully immersed during washes and antibody incubations.

Contaminated equipment. Protein or pieces of gel remaining on the unit may stick to the membrane. Antibody can get trapped in the gel, and then are washed out poorly, resulting in intense localized signal.

Aggregation of blocking agent. When blocking agent is powder stir it over night at 4°C to make sure it is completely dissolved.

Interaction of the membrane with sample tray. Always use clean plastic trays to avoid any type of cross-reaction.

Formation of aggregates in HRP-conjugate. Filter secondary antibody solution through a 0.2 μm filter. Use fresh antibody.
No bands or weak bands

**Excessive signal generated.** The enzyme in the system depleted the substrate and caused the signal to fade quickly. Further dilute secondary Ab.

**Inefficient transfer.** Ensure that there is good contact between membrane and gel during sandwich assembling. High MW protein may require more time for transfer. Reduce voltage or time of transfer for low molecular weight proteins (< 10 kDa).

**Antibodies may have lost activity.** Perform a Dot Blot. Follow manufacturer's recommended storage and avoid freeze/thaw cycles.

**Incorrect secondary antibody used.** Confirm host species/Ig type of primary Ab.

**Low protein-antibody binding.** Reduce the number of washes to minimum. Reduce NaCl in TBS-T Buffer (100mM-350mM).

**Non-fat dry milk may mask some antigens.** Decrease blocking time. Decrease milk percentage in Blocking Buffer or substitute with 5% BSA Blocking Buffer.

**Sodium azide contamination.** Make sure buffers do not contain sodium azide as this will quench HRP signal.

**Contaminated stock solutions.** Do not contaminate the chemiluminescent substrate stock solutions using the same pipette tip. Use new reagents.

Non-specific bands

**Aggregation of analyte.** Increase amount of reducing agent to ensure complete reducing of disulfide bonds.

**SDS interference.** The presence of SDS may result in the development of unspecific bands caused by antibodies binding to the charged SDS molecules associated with the proteins. Wash thoroughly the membrane after transfer with water.

**High protein concentration.** A commonly seem effect is the diffusion of protein bands. Reduce the amount of protein initially loaded.

**Primary antibody is not specific for the protein of interest.** Use monospecific or antigen affinity purified antibodies. Always incubate your primary antibody at 4°C overnight and not at room temperature. Reduce NaCl in TBS-T Buffer (100mM-350mM). Use monospecific or antigen affinity purified Ab.

**Non-specific binding of secondary antibody.** Confirm the secondary is specific by omitting the primary and running a secondary only blot. If bands develop choose an alternative secondary antibody.
**White bands or “ghost bands”**

**Excessive signal generated.** Excessive antibodies or loaded protein can cause high levels of localized signal. This results in rapid consumption of substrate at this point. Since there is no light production after the completion of this reaction, white bands are the result. Try first to further dilute secondary antibody.

**Uneven or jagged bands**

**Uneven gel run.** Load all available wells. Empty wells can be loaded with sample buffer.  
**Voltage or current were too high during electrophoresis.** Reduce voltage or current during electrophoresis.

**Effects of high salt in samples.** Reduce NaCl concentration in TBS-T Buffer (100mM÷350mM).
### 11. Selection guide

<table>
<thead>
<tr>
<th>Product</th>
<th>Competitors</th>
</tr>
</thead>
<tbody>
<tr>
<td>WESTAR SUN</td>
<td>AMERSHAM™ ECL™ - GEHC AMERSHAM™ ECL™ START - GEHC PIERCE™ECL - THERMO SCIENTIFIC™ LUMINATA™ CLASSICO - MILLIPORE™</td>
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<tr>
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<tr>
<td>WESTAR NOVA 2.0</td>
<td>SUPERSIGNAL™ WEST PICO - THERMO SCIENTIFIC™ LITEABLOT® PLUS - EUROCLONE LUMINATA™ CLASSICO - MILLIPORE™ WESTERN LIGHTNING™ PLUS - PERKINELMER</td>
</tr>
<tr>
<td>cod. XLS071,0250</td>
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</tr>
<tr>
<td>WESTAR ETA C 2.0</td>
<td>CLARITY™ - BIO-RAD PIERCE™ECL™ PLUS - THERMO SCIENTIFIC™ LITEABLOT® EXTEND - EUROCLONE SUPERSIGNAL™ WEST DURA - THERMO SCIENTIFIC™ LUMINATA™ CRESCEndo - MILLIPORE™</td>
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<tr>
<td>WESTAR ETA C ULTRA 2.0</td>
<td>ECL™ PRIME - GEHC SUPERSIGNAL™ WEST DURA - THERMO SCIENTIFIC™ LITEABLOT® EXTEND - EUROCLONE LUMINATA™ FORTE - MILLIPORE™ IMMUBILON™ - MILLIPORE™ WESTERN LIGHTNING PRO - PERKINELMER</td>
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<tr>
<td>WESTAR SUPERNOVA</td>
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</table>

For further information, visit [www.cyanagen.com](http://www.cyanagen.com)

For orders: call +39 051.534063
mail to sales@cyanagen.com