

WESTERNLUMAXLIGHT

310208	WesternLumaxLight ^{1M} Superior
310209	$We stern Lumax Light {}^{TM}Superior$
310212	$We stern Lumax Light^{TM} Enhance$
310231	WesternLumaxLight TM Sirius

⁻More sensitive signal than Thermo ECL or Pierce ™Super signal ECL.

⁻Perfect performance for x-ray film imaging and CCD imaging.

⁻Detecting petagram protein amounts.

⁻Signal will be last to 2 hours.

WesternLumaxLightTM

Protocol

- 1. Prepare your protein blot on either PVDF or nitrocellulose using your standard technique.
- 2. Block membrane for one hour at room temperature(RT).
- 3. Incubate blot with primary antibody for one hour at RT with gentle agitation.
- 4. Wash blot with PBS-T or TBS-T:
 - 1× quickly.
 - 1×15min, with 0.7ml/cm2 membrane each time.
- 5.incubate blot with secondary antibody for one hour at RT with gentle agitation.
- 6. Wash blot with PBS-T or PBS-T:
 - 3×5min, with an least 0.5ml/cm2 membrane each time.
- 7. Mix WesternLumaxLight components 1:1 in sufficient amounts to obtain at least 0.1cm/ml2 of your membrane and place on blot for 2 minutes.

(For example, for a 7×9cm blot, mix at least 3.15ml of each component to obtain 6.3ml working reagent.)

- 8. Remove the membrane from the reagent and place it on the provided Background quenching sheet. Drain excess reagent by holding the sheet with the membrane vertically for a few moments.
- 9. Place the plastic sheet with the blot in your CCD imager and image.

If a very long exposure is required to detect weak bands, or if imaging will be done using X-ray film, cover the damp blot with plastic wrap.