

# **EZ-ECL**

**Chemiluminescence  
Detection Kit for HRP**

Cat. No.: 20-500-120



BIOLOGICAL INDUSTRIES  
ISRAEL, BEIT HAEMEK LTD.



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## **BIOLOGICAL INDUSTRIES EZ-ECL**

### Protocol for Proteins

Step	Action	Volume	Time	Remarks
Electrophoresis and Blotting	According to the usual protocols			Use Nitrocellulose or PVDF membrane
Membrane Blocking	Block membrane with blocking solution, TBS-T or PBS-T with 5% dried milk (w/v), under constant shaking	0.5ml/cm <sup>2</sup>	1 hour at room temperature	Alternatively: overnight at 4°C without shaking
Primary Antibody	Dilute the primary antibody in TBS-T or PBS-T with 2% dried milk (w/v). Incubate the membrane in solution with shaking	0.1ml/cm <sup>2</sup>	1 hour at room temperature	Alternatively: overnight at 4°C without shaking
Washing	Three times with TBS-T under constant shaking	0.5ml/cm <sup>2</sup>	3 x 10 minutes	
Secondary Antibody	Dilute the HRP-labeled secondary antibody (1:10,000-1:60,000) in TBS-T or PBS-T with 2% dried milk (w/v). Incubate the membrane in the solution	0.1ml/cm <sup>2</sup>	1 hour at room temperature	
Washing	Three times with TBS-T under constant shaking	0.5ml/cm <sup>2</sup>	3 x 10 minutes	
Equilibration	Mix equal volumes of EZ-ECL solution A&B	0.1ml/cm <sup>2</sup>	5 minutes	
Detection	Incubate membrane in detection mix solution	0.1ml/cm <sup>2</sup>	1-3 minutes	With gentle shaking
Exposure	Remove excess detection mix, wrap in saran wrap and expose to film		0.5-60 minutes	Remove air pockets

## EZ-ECL Chemiluminescence Detection Kit for HRP

Cat. No.: 20-500-120 Store at: 24°C

**Product Description**  
 EZ-ECL is a complete kit with ready-to-use reagents for chemiluminescent detection of immobilized proteins (Western blotting or immunodiffusion) or nucleic acids (Southern or Northern), conjugated with HRP directly or indirectly. The use of enhanced chemiluminescence was introduced by Thorpe and Kricka (1,2). In the presence of hydrogen peroxide ( $H_2O_2$ ), horseradish peroxidase (HRP) catalyzes the oxidation of cyclic dihydrazides, such as luminol. Immediately following the oxidation, the luminol is in an excited state (intermediate reaction product), which decays to the ground state by emitting light. Strong enhancement of the light emission is produced by enhancers such as phenolic compounds. Using this method, it is possible to detect membrane immobilized specific antigens, or sequences of nucleic acids, labeled directly with HRP or indirectly with HRP-labeled antibodies/streptavidin.

### Principles of Protein Detection Procedure

Advantages of using the EZ-ECL Detection Kit  
 - High sensitivity non-radioactive detection system.  
 - Stable hard copy results on film.  
 - Only small amounts of antibody required.  
 - Detection may be achieved in short exposure times (minutes)  
 - High resolution.

Kit Reagents  
 Cat. No.: 20-500-120A EZ-ECL Solution A, 60ml  
 Contains: Luminol and enhancer  
 Store at: 2-8°C.

Caution  
 If above solutions come into contact with eyes or skin, flush with plenty of water and remove contaminated clothing.

## Protocol for Western Blotting and Chemiluminescence Detection

### 1. Preparation of Solutions

- 1.1 Tris Buffered Saline (TBS)  
 6.03g Tris base (50mM)  
 8.76g Sodium Chloride (150mM)  
 Adjust pH to 7.5 with Hydrochloric Acid  
 Add distilled water up to 1000ml
- 1.2 Phosphate Buffered Saline (PBS) - optional  
 11.3g Di-sodium Hydrogen Phosphate, anhydrous (80mM)  
 2.96g Sodium Dihydrogen Phosphate (20mM)  
 5.84g Sodium Chloride (100mM)  
 Add distilled water up to 1000ml  
 Check pH (should be 7.5)
- 1.3 TBS-Tween (TBS-T) and PBS-Tween (PBS-T)  
 Dilute 1ml of Tween-20 in 1000ml of buffer (0.1% final concentration).
- 1.4 A sufficient volume of wash buffer, blocking buffer, and antibody solution should be used to cover the blot to ensure that the membrane does not become dry. This will also ensure a reduced non-specific background.
- 1.5 Do not use sodium azide as a preservative for the secondary antibody dilutions, as azide irreversibly inhibits horseradish peroxidase.
- 1.6 Wherever use of dried milk is indicated, this can be substituted with low-fat milk.

### 2. Electrophoresis, Blotting and Membrane Preparation

- 2.1 Carry out electrophoresis for protein separation. Either non-denaturing gel SDS-PAGE or two-dimensional gels may be used.
- 2.2 Transfer proteins from the gel to a membrane. Use nitrocellulose or PVDF membrane. PVDF membranes must be wetted briefly in methanol then soaked in distilled water for 1-3 minutes, followed by equilibration in transfer buffer.
- 2.3 Membrane Blocking  
 Block non-specific binding sites by incubating the membrane for 1 hour at room temperature with shaking in TBS-T or PBS-T solution containing 5% dried milk (w/v). This step can be performed overnight at 4°C without shaking.
- 2.4 Primary Antibody  
 Dilute the primary antibody in TBS-T or PBS-T with 2% dried milk (w/v). Incubate the membrane in the solution for 1 hour at room temperature with shaking, or overnight at 4°C, without shaking.
- 2.5 Membrane Washing  
 Wash the membrane three times in TBS-T or PBS-T for 10 minutes each. Use at least 50ml of buffer for 10cm membrane.

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### Protocol for Nucleic Acids

Step	Action	Volume	Time	Remarks
Electrophoresis	According to the usual protocol			
Blotting	Immerse the membrane in ddH2O and then in transfer buffer. Transfer according to the usual protocol			
Membrane Washing and Fixation	- soak the membrane in 6xSSC - dry for at least 30 minutes - fixation by baking or by U.V. crosslinking	0.5ml/cm <sup>2</sup>	- 5 minutes at room temperature - room temperature - 0.5-2 hours at 80°C or 32 seconds 1200 erg.	Use Nitrocellulose or nylon membrane
Pre-hybridization and Hybridization	According to the manufacturer's instructions Use denatured biotin-labeled DNA probe	0.1ml/cm <sup>2</sup>	1-2.5 hours at 68°C or 60°C	
Washing (I)	Twice with 2xSSC, 0.1% SDS (Wash No. 1)	0.5ml/cm <sup>2</sup>	2 x 15 minutes at room temperature	
Washing (II)	Twice with 1-0.1xSSC, 0.1% SDS (Wash No. 2)	0.5ml/cm <sup>2</sup>	2 x 15 minutes at 68°C or 60°C	
Blocking	Immerse the membrane in Buffer A Incubate the blot in 0.2% EZ-Block in Buffer A	2ml/cm <sup>2</sup> 2ml/cm <sup>2</sup>	30 minutes at room temperature	
Streptavidin-HRP	Dilute the streptavidin-HRP (1:2000-1:10,000) in 0.5% EZ-Block in Buffer A. Incubate the membrane in the solution.	0.125ml/cm <sup>2</sup>	30 minutes at room temperature	
Washing	Three times with 0.1% Tween 20 in Buffer A	2ml/cm <sup>2</sup>	3 x 10 minutes	
Equilibration	Mix equal volumes of EZ-ECL solution A&B	0.1ml/cm <sup>2</sup>	5 minutes	
Detection	Incubate membrane in detection mix solution	0.1ml/cm <sup>2</sup>	1-3 minutes	With gentle shaking
Exposure	Remove excess detection mix, wrap in saran wrap and expose to film	0.5-60 minutes		Remove air pockets

## EZ-ECL

Problem	Possible Cause	Suggested Solution
Large "Blothy" Areas of Background	Undiluted probe connected to the membrane. Improper blocking	Thoroughly mix the probe, diluted in hybridization solution before pouring it onto the membrane. <ul style="list-style-type: none"> <li>• Check that blocking agent solution has been made properly.</li> <li>• Use a fresh prepared solution of blocking agent.</li> <li>• Increase concentration of blocking agent.</li> <li>• Include blocking agent in HRP conjugate solution.</li> <li>• Increase Tween 20 concentration of blocking solution.</li> <li>• Increase incubation time and/or temperature of blocking incubation.</li> <li>• Use hemispherical blocking agents freshly prepared.</li> <li>• 1-10% bovine serum albumin in TBS-T or PBS-T.</li> <li>• 0.5-3% gelatin in TBS-T or PBS-T.</li> <li>• 1% polyvinyl pyrrolidone (PVP) in TBS-T or PBS-T.</li> </ul>
Improper washing	Problems with membranes Improper washing times and volumes of wash buffers Improper washing	<ul style="list-style-type: none"> <li>• Add Tween 20 to solution if not already included in the solution.</li> <li>• Increase wash stringency if needed.</li> <li>• Check that membranes are completely rinsed in all solutions, especially during washing.</li> <li>• Use a fresh supply of membranes.</li> <li>• The type of membrane used was not compatible with non-radioactive systems.</li> <li>• Hatch blots carefully with forces and blunt non-serIALIZED forces. Damage to the membrane can cause non-specific binding of the probe or the detection reagents.</li> <li>• Use clean forceps to handle blots after washing.</li> </ul>
Problems with membranes	HRP-conjugate concentration is too high Detection reagents	<ul style="list-style-type: none"> <li>• Optimize HRP-conjugate concentration to reduce the background.</li> <li>• Rewash blot twice for 10 minutes in wash buffer, and repeat detection steps.</li> <li>• Assess detection reagents in blots. Dark wells by absorbing the excess detection reagent before placing the blots in film cassettes.</li> </ul>
Overexposure	Overexposure	<ul style="list-style-type: none"> <li>• Expose film for minimum period 5-10 seconds.</li> <li>• Leave blot in the cassette for 5-10 minutes before re-exposing to film.</li> </ul>
Contaminated buffers	Contaminated blotting equipment	<ul style="list-style-type: none"> <li>• Use fresh buffers.</li> <li>• Clean or replace equipment.</li> </ul>
Uneven Spotted Blot	Improper blotting technique Unevenly hydrated membrane	<ul style="list-style-type: none"> <li>• Check that gel and membrane make proper contact during blotting.</li> <li>• Use new membranes.</li> <li>• Make sure that membrane is fully covered and weighed during incubations.</li> </ul>
	HRP conjugate solution is not completely clear	<ul style="list-style-type: none"> <li>• Centrifuge the HRP-conjugate solution and use supernatant before diluting in blocking solution. Alternatively, filter through a 0.2-μm filter with low protein absorption.</li> </ul>
	Fingerprints and/or keratin contamination	<ul style="list-style-type: none"> <li>• Avoid touching membrane. Use gloves and blunt forceps.</li> </ul>

Problem	Possible Cause	Suggested Solution
<b>Excessive Diffuse Signal</b>	Overloading of protein Improper gel conditions Increase arylamine concentration of gel. Check gel and buffer recipes. Check that no bubbles remaine with transfer from gel to membrane.	Dilute the HRP labeled secondary antibody in TBS-T or PBS-T with 2% dried milk (w/v). Incubate the membrane in the solution for 1 hour at room temperature with shaking. Wash the membrane as detailed in 2.5.
<b>For Nucleic Acid Blots</b>	No, or inefficient transfer Or Weak Signal	2.6 Secondary Antibody Dilute the HRP labeled secondary antibody in TBS-T or PBS-T solution containing 5% dried milk (w/v). This step can be performed overnight at 4°C without shaking. 2.7 Membrane Washing Wash the membrane as detailed in 2.5.
<b>High Background</b>	Film exposure time is too short Insufficient pre-hybridization and/or blocking	4.1.3 Block non-specific binding sites by incubating the strip for 1 hour at room temperature with shaking in TBS-T or PBS-T solution containing 5% dried milk (w/v). 4.1.4 Prepare several dilutions of primary antibody in TBS-T or PBS-T with 2% dried milk (w/v), e.g. 1:100-1:5,000. Incubate one piece of membrane in each dilution for 1 hour at room temperature with constant shaking, or overnight at 4°C without shaking. 4.1.5 Wash the membranes three times in TBS-T or PBS-T for 10 minutes each. Use at least 0.5 ml of buffer per 1cm <sup>2</sup> membrane. 4.1.6 Dilute the HRP-labeled secondary antibody in TBS-T or PBS-T with 2% dried milk (w/v) to the known optimal dilution. Incubate each strip in the solution for 1 hour at room temperature with shaking. 4.1.7 Wash the membranes as detailed in 4.1.5 above. 4.1.8 Detection: as detailed in 2. above. 4.2 Dot-Blot for Secondary Antibody Optimization Prepare one piece of nitrocellulose membrane for each secondary antibody to be tested. 4.2.1 Pre-dilutions as detailed in 4.1.1 - 4.1.3 above. 4.2.2 Dilute the primary antibody in TBS-T or PBS-T with 2% dried milk (w/v) to the known optimal dilution. Incubate each strip in the solution for 1 hour at room temperature with shaking. 4.2.3 Wash the membranes as detailed in 4.1.5 above. 4.2.4 Prepare several dilutions of secondary antibody in TBS-T or PBS-T with 2% dried milk (w/v), e.g. 1:5,000-1:100,000. Incubate one piece of membrane in each dilution for 1 hour at room temperature with constant shaking. 4.2.5 Wash the membranes as detailed in 4.1.5 above. 4.2.6 Detection: as detailed in 2. above.

Problem	Possible Cause	Suggested Solution
<b>No Signal</b>	Overloading of protein Improper gel conditions Increase probe concentration during hybridization, and/or expose blot to film for longer period.	3.1 Preparations Prepare the following equipment and solutions in a dark room: -X-ray film cassette -X-ray film -Tinner -Developer, fixer and water in tanks -Transparent plastic bag or saran wrap -Glass pipettes -Sterile gloves - to prevent hand contact with membrane, film or reagents
<b>3. Enhanced Chemiluminescence Detection</b>	Probe not completely on or for 5 minutes.	3.2 Detection 3.2.1 Mix an equal volume of EZ-ECL Solution A and EZ-ECL Solution B to give sufficient solution to cover the membrane (0.1ml/cm <sup>2</sup> ). Let the detection mix equilibrate for at least 5 minutes. 3.2.2 Drain the excess buffer from the washed blots. Do not let the membrane dry out. Add the detection mix directly to the blot (protein side up) Incubate for 1-3 minutes at room temperature. 3.2.3 Drain off excess detection mix and wrap the membrane in saran wrap. 3.2.4 Place the blot(s) Protein side up, in the film cassette. Switch off the lights and use red safety light. Place a sheet of film on the blot, close the cassette and expose for 30-60 seconds. 3.2.5 Replace the exposed film with a new one, close the cassette and develop the first exposed film. 3.2.6 Expose the second film for a suitable time according to the signal intensity on the first film. 3.2.7 If signal intensity was too high, wait up to 30 minutes before re-exposing.
<b>4. Optimization of Antibody Concentration for EZ-ECL</b>	The immunoblot can be stripped off blocking reagent and antibodies, and then reprobed as required.	5. Stripping and Reprobing of Membrane It is essential to optimize the immunoblot conditions to achieve maximum signal and minimum background. First optimize the concentration of the primary antibody using a constant amount of secondary-HRP conjugate. Using the optimized primary antibody concentration, adjust the concentration of the secondary antibody-HRP conjugate.
<b>High Background</b>	Film exposure time was too high during hybridization Insufficient pre-hybridization and/or blocking	5.1 Incubate membrane in stripping buffer for 30 minutes at 50-70°C. (62.5 mM Tris-HCl pH 6.8, 100mM β-mercaptoethanol and 2% (w/v) SDS). 5.2 Wash the membrane twice in TBS-T or PBS-T for 10 minutes each. Use at least 50 ml of buffer for 10cm <sup>2</sup> membrane. To ensure removal of antibodies, include a piece of nitrocellulose membrane for each primary antibody dilution to be tested. 4.1.1 Spot a dilution range of protein onto the membrane. 4.1.2 Allow the membrane to air-dry. 5.3 Reprobe the blot as detailed in 2.3.-3.2.7 above.

Problem	Possible Cause	Suggested Solution
<b>High Background (cont.)</b>	Inappropriate blocking (cont.)	• Increase Tween 20 concentration (Tween 20 may reduce the binding of antibodies, especially if low affinity primary antibodies). • Increase incubation time and/or temperature of blocking incubation. • Try alternative blocking agents freshly prepared: -1-10% bovine serum albumin in TBS-T or PBS-T. -0.5-2% gelatin in TBS-T or PBS-T. -1% bovine pyromelitone (PVP) in TBS-T or PBS-T.
<b>Problems with membranes</b>	Improper washing	• Increase washing times and volumes of wash buffers. • Add Tween 10 to solutions if not already present. • Check that membranes are completely immersed in all solutions, especially during washing. • Use a fresh supply of membranes. • Handle blots carefully with gloves and blunt non-sterile forceps. Damage to the membrane can cause non-specific binding of the detection reagents. • Use clean forceps to handle blots after washing.
<b>Antibody concentration is too high</b>	Optimize antibody concentration to reduce the background.	• Rewash blots twice for 10 minutes in wash buffer and repeat detection steps. • Excess detection reagents in blots. Draw well by absorbing the excess on tissue paper before placing the blots in film cassettes.
<b>Detection reagents</b>	Overexposure	• Expose the film for a minimum period (5-30 seconds). • Leave blots in the cassette for 5-10 minutes before re-exposing to film.
<b>Contaminated buffers</b>	Contaminated blotting equipment	• Use fresh buffers. • Clean or replace equipment.
<b>Unevenly hydrated membrane</b>	Inappropriate blotting technique	• Check that gel and membrane make proper contact during blotting. • Check that excess temperatures are not reached during electroblotting, producing bubbles, gel/membrane distortion, etc. • Use new membranes. • Make sure that membrane is fully covered and wicked during incubations.
<b>Primary antibody and/or HRP conjugate solutions are not completely clear</b>	Fingerprints and/or keratin contamination	• Re-purify the antibodies and use supernatant before diluting in blocking solution. Alternatively, filter through a 0.2μm filter with low protein absorption. • Avoid touching membrane. Use gloves and blunt forceps.

## **E-Z-*FCI* Troubleshooting Guide For Protein Blots**

### **Protocol for Southern/Northern Blotting and Chemiluminescence Detection**

Problem	Possible Cause	Suggested Solution
No Signal Or Weak Signal	No or inefficient protein transfer.	Transfer conditions were improper. Check that gel and blotting membrane are correctly oriented with the apole side facing the transfer, stain gel and membrane to check transfer efficiency.
	Insufficient protein was loaded onto the gel.	Increase amount of protein applied to the gel.
	Protein degradation on membrane stored before detection.	<ul style="list-style-type: none"> <li>Check storage conditions of membrane; target protein degradation may occur if the blots are stored incorrectly.</li> <li>Use fresh blots.</li> </ul>
	Primary antibody does not detect denatured proteins (native protein in plaque) if the primary antibody only binds to native protein. Try to use a non-denaturing SDS or urea.	Perform dot blot with denatured protein and native protein in plaque if the primary antibody only binds to native protein.
	Affinity of primary antibody is low.	<ul style="list-style-type: none"> <li>Optimize antibody concentration.</li> <li>Prolong incubation with primary antibody to overnight at 4°C.</li> <li>Shorten washing times and use washing buffer without Tween 20.</li> <li>Increase protein antibody in buffer without blocking reagent. If staining may be increased, prolong detection time.</li> </ul>
	Concentration of secondary antibody is too low.	<ul style="list-style-type: none"> <li>Dot different dilutions of HRP-conjugate onto membrane and detect directly. If no signal appears, use fresh HRP-conjugate and test in the same way. If still no signal appears, check E-Z-<i>FCI</i> detection reagents.</li> </ul>
	E-Z- <i>FCI</i> detection reagent gives no signal.	<ul style="list-style-type: none"> <li>E-Z-<i>FCI</i> detection reagents may have become contaminated.</li> <li>Incorrect storage of the E-Z-<i>FCI</i> reagents may cause loss of signal.</li> </ul>
	Film exposure time is too short.	Expose film for extended period (1-2 hours).
High Background	Improper blocking.	<p><u>Notes:</u></p> <ul style="list-style-type: none"> <li>Nylon membrane binds small DNA fragments more efficiently than nitrocellulose membranes.</li> <li>Fragments of less than 300 nucleotides in length are not retained by 0.45 micron nitrocellulose membranes. Use a pore size of 0.2 micron.</li> <li>Use gloves and blunt-ended forceps to handle the membrane.</li> </ul> <p><b>2.4 Soak the nitrocellulose membrane in de-ionized water until completely wet.</b></p> <p><b>2.5 Immerse the membrane in transfer buffer (20×SSC, or 20×SSPE).</b></p> <p><b>2.6 Transfer the nuclear acids from the gel to a membrane for 2-24 hours. Mark the positions of the gel slots on the filter with a very soft lead pencil or a ball point pen.</b></p> <p><b>2.7 Remove the membrane from the 20×SSC and allow excess fluid to drain away. Place the membrane flat on a paper towel to dry for at least 30 minutes at room temperature.</b></p> <p><b>2.8 Sandwich the filter between two Sheets of dry 3MM Paper. Fix the DNA to the filter by baking for 30 minutes to 2 hours at 80°C in a vacuum oven.</b></p> <p><b>2.9 Hybridization using EZ-Hybridization Solution with non-radioactively labeled probes.</b></p> <p><b>2.10 Wash the EZ-Hybridization Solution at 68°C for Northern and at 60°C for Southern, and stir well to completely dissolve any precipitate.</b></p> <p><b>2.11 Pour hybridization mixture in a minimum of 0.1 ml/cm<sup>2</sup> of EZ-Hybridization mixture over the filter.</b></p> <p><b>2.12 Seal the filter with Parafilm® and incubate at 68°C for Northern and at 60°C for Southern for 30-90 minutes. The volume of solution must be sufficient to completely cover the membrane, or high backgrounds may result.</b></p> <p><b>2.13 Denature the non-radioactively labeled DNA probe at 95-100°C for 2-5 minutes. Chill quickly on ice.</b></p> <p><b>2.14 Add non-radioactively labeled probe to a sufficient volume of fresh EZ-Hybridization Solution. Mix gently. For recommended final probe concentrations, see notes above.</b></p> <p><b>2.15 Replace the EZ-Hybridization Solution with the fresh solution containing the non-radioactively labeled DNA probe. Remove all air bubbles from the container and make sure the EZ-Hybridization Solution is evenly distributed over the entire blot.</b></p> <p><b>2.16 Incubate the filter with the hybridization mixture at 68°C for Northern and at 60°C for Southern for 1-2.5 hours. (For high target applications, shorter hybridization times can be used. For single gene sequences, hybridization can be performed overnight).</b></p> <p><b>2.17 Wash the membranes at room temperature twice, 15 minutes each time, with at least 0.2 ml/cm<sup>2</sup> of 2×SSC, 0.1% SDS (Wash No. 1).</b></p> <p><b>2.18 Wash the membrane twice at 68°C for Northern and at 60°C for Southern, 15 minutes each time, with at least 0.5 ml/cm<sup>2</sup> of 1-1×SSC, 0.1% SDS, with continuous agitation.</b></p> <p><b>2.19 Wash the membrane twice at 68°C for Northern and at 60°C for Southern, 15 minutes each time, with at least 0.5 ml/cm<sup>2</sup> of 1-1×SSC, 0.1% SDS, with continuous agitation.</b></p> <p><b>2.20 Expose the film for a suitable time according to the signal intensity on the first film.</b></p> <p><b>2.21 If signal intensity was too high, wait up to 30 minutes before re-exposing.</b></p> <p><b>2.22 Denature the DNA by soaking the gel for 45 minutes in several volumes of 1.5M NaCl, 0.5M NaOH with constant, gentle agitation.</b></p> <p><b>2.23 Rinse the gel briefly in de-ionized water, and neutralize it by soaking for 30 minutes in several volumes of a solution of 1M Tris pH 7.4, 1.5M NaCl at room temperature with constant, gentle agitation. Change the neutralization solution and continue soaking the gel for a further 15 minutes.</b></p> <p><b>2.24 Check that blocking agent solution has been made properly.</b></p> <p><b>2.25 Use a freshly prepared solution of blocking agent.</b></p> <p><b>2.26 Increase concentration of blocking agent.</b></p> <p><b>2.27 Include blocking agent in antibody solutions.</b></p>

- Notes:**
- (1) Thorne, C.H.C., and Kricka, L.J., Methods in Enzymology, 133:331-353 (1986)
  - (2) Thorne, C.H.C., Kricka, L.J., Moseley, S.B. and Whitehead, T.P., Clin. Chem., 31(8):1335-1341 (1985)
  - (3) Riko, I. et al, Analytical Biochemistry, 231:170-174 (1995)

- References:**
- (1) Thorne, C.H.C., and Kricka, L.J., Methods in Enzymology, 133:331-353 (1986)
  - (2) Thorne, C.H.C., Kricka, L.J., Moseley, S.B. and Whitehead, T.P., Clin. Chem., 31(8):1335-1341 (1985)
  - (3) Riko, I. et al, Analytical Biochemistry, 231:170-174 (1995)

- Note:**
- These washing conditions may be too stringent for probes that are not completely homologous to the target. If this is the case, lower the temperature to 50°C.