## **General Protocol for Western Blot Development**

**Pre-preparation**: Make available target proteins from cell and tissue lysates separated by SDS PAGE and blotted onto PVDF membrane.

- 1. Cut and label required strips (35ug protein each)
- 2. Soak the strips in methanol for 10 seconds.
- 3. Wash the strips first in TBS for 5 minute, then in TBST for 15 minute. Except for the incubation in methanol, all other incubations should be carried out on a rocking table.
- 4. Block the strips by blocking buffer (3% skimmed milk in TBST) for 40 minutes.

## **Development and Detection of the Primary antibodies**

- 5. Incubate the strips with primary antibody (diluted e.g. lug/ml in blocking buffer -please refer to datasheet for recommended concentration) for 1 hour at room temperature. Please note: Primary incubation overnight at 4°C requires significant dilution of the primary antibody and non-specific background may result.
- 6. Wash the strips with TBST for 3 x 10minutes.
- 7. Incubate the strips with diluted secondary antibody (1:40K) in blocking buffer for 1 hour.
- 8. Wash the strips for 3 x 10minutes.
- 9. Incubate strips with chemiluminescence substrate (100ul/strip) for 3 minutes.
- 10. Develop the bands on x-ray film in dark room under the red light for approx 15 seconds or determine based upon signal in films.
- 11. Label the detail on the film including molecular weight marking of strips.
- 12. Scan the film and read the molecular size and strength of bands.

## Further Guidelines:

- Tris buffered saline (TBS): 20mM Tris, pH7.4 in 150mM NaCl
- Include a negative control lane this may be a lysate, or just omission of the Primary antibody.
- Where possible include a positive control lysate (e.g.as per datasheet)