

## Principle and Methodology

**Hellabio Immunofixation Electrophoresis (IFE) kits** are intended for *in vitro* diagnosis of monoclonal paraproteins in human serum and other biological samples.

In some pathological cases (multiple myeloma etc) abnormal monoclonal bands appear in electropherogram. The identification of these monoclonal bands can be done by different immunological techniques such as Immunoelectrophoresis and Immunofixation electrophoresis (IFE).

The IFE technique combines zone electrophoresis with immunoprecipitation and is easier and more practical than other techniques. By IFE proteins of sample are first according to their charge separated by electrophoresis on the agarose film, and then they (as antigen) are reacted and fixed with monospecific polyclonal antisera. The monoclonal paraproteins can be heavy chain of IgG, IgA, IgM, (IgD, IgE) and or light chain of kappa or lambda.

Hellabio antisera kit contains Protein fixation solution, the anti heavy chain of IgG, IgA, IgM, and free+bound anti light kappa and lambda chain.

**If specific antigen (heavy and or light chain globulins) is present, a characteristic immunoprecipitin band will be formed, which is visualized by staining of the gel with Amido Black.**

**Required Reagents and Equipment's** included in each kit:  
[Warning: All reagents from each kit must be used together]

Product	KIT IFEAD01 /20 TESTS
Agarose Gels	10
Sponges	20
Buffer Solution [3X concentrated]	50 ml
Staining Solution [5X concentrated]	60 ml
Diluent solution [ready to use]	12 ml
Destaining solution [500X concentrated]	20 ml
Sample Templates	10
Antisera templates	10
Drying Blotter Sheets	50
Gel Blotter Sheets	60
Antisera [ Anti IgG ( $\gamma$ ), IgA ( $\alpha$ ), IgM ( $\mu$ ), Goat anti-Human antisera (f+b) $\kappa$ - and $\lambda$ -chain ]	
Protein fixation solution	1.5 ml
Instructions for use in English	

**All reagents must be used according to the instructions and until the expiration date indicated on the kit**

### Preparation, storage and stability of the reagents included in the kit:

**a) Agarose Gels:** Agarose Gels are in non- barbital buffer and other non-reactive ingredients for long stability and optimum resolution of protein fractions. The Gels must be stored at **15 - 25 °C on horizontal position** until the expiration date indicated on the kit. Do not freeze the gels. Carefully discover the gel just before use and follow the instructions of the manual.

**b) Diluent Solution:** Working electrophoresis buffer + Bromphenol blue and other non-reactive ingredients. Store it at 15 - 25 °C until the expiration date indicated on the kit. **Ready to use.**

**c) Electrophoresis Buffer for sponges: 3X concentrated** stock solution of non barbital buffer and important non-reactive ingredients. To prepare working solution dilute it 1:3 [dilute the buffer to 100ml distilled water]. To use put the sponges on the plastic plate on horizontal position and add to each sponge 6.5 ml of working buffer solution. The working buffer solution can be stored at 4-8 °C until the expiration date indicated on the kit.

**d) Staining Solution:** Concentrated Amido Black solution. Store the concentrated solution at 15 - 25 °C until the expiration date indicated on the kit. To prepare working solution dilute the content of the bottle according to the instructions on the bottle. The diluted solution is enough for the staining of all gels of the kit. It should be stored in a closed flask at room temperature until 3 months.

**e) Gel Blotter strips:** Thin filter paper strips to blot the gel in the application area. Avoid humidity. Blot just for 5 seconds. Avoid humidity.

**f) Gel Blotter Sheets:** Thin filter paper sheet to blot the entire surface of the gel.

**g) Drying Blotter Sheets:** Thick filter paper sheet which absorbs the excess humidity of the gel.

**h) Destaining solution:** 2% citric acid solution. To prepare working solution dilute the content of the bottle according to the instructions on the bottle. Store at room temperature until the expiration date indicated on the kit.

**i) Antisera kit:** Each kit contains protein fixation solution, goat anti-Human heavy chain of IgG, IgA and IgM and free+bound anti light chain of kappa ( $\kappa$ ) and lambda ( $\lambda$ ). Store at -18 °C (long term condition) until the expiration date indicated on the kit.

### Limitation / Caution:

- Do not use the Agarose gel if it seems to be dried or infected by fungi.
- Do not freeze the agarose gel.
- Store the kit in horizontal position.
- Do not use hemolytic sera or plasma.

### Collection and handling of specimens:

The samples should be collected according to the standard hospital methods.

**a) Blood:** Analysis is preferably performed on sera in order to avoid the fibrinogen band (in plasma), which migrates in the gamma zone and could lead to a false interpretation. Venous blood is drawn into tube. After centrifugation the supernatant fluid (serum) can be stored at 4-8°C for 48-72 hours. Storage at -20°C may disturb some of the electrophoretic patterns. Care should be taken to prevent haemolysis in the serum because it will cause false elevation in the  $\alpha_2$  and  $\beta$ -fractions. Samples with much precipitate that will not dissolve on warming to 37°C should be either centrifuged or allowed to settle before the clear supernatant fluid is applied to the gel surface.

**b) Cerebrospinal fluid:** Normally, some 80% of the proteins of cerebrospinal fluid originate from plasma, and the rest have been synthesised locally. The filtration resistance for plasma protein rapidly increases with molecular size and therefore the ratio of large proteins normally is much lower in CSF than in plasma. The protein content of CSF is in the range of 15-40 mg/dl. Therefore it must be concentrated at least 150-fold before electrophoresis.

**c) Urine:** Analysis of the urinary proteins shows that normal urine contains more kind of proteins which are much diluted and therefore it must be concentrated. For the detection of Bence-Jones proteins, free Kappa and Lambda light chains, the sample has to be concentrated to  $\geq$  100 mg/dl of total protein and to about 80 mg/dl of total proteins for immunoglobulin.

## Procedure of Immunofixation electrophoresis

Extreme care must be taken in choosing the appropriate dilution of the serum sample (100 – 200 mg/dl). The incorrect choice of serum dilution may result in either inability to detect a minor monoclonal protein or a prozone effect. When the protein concentration is low (cerebrospinal fluid, urine), it must be concentrated to get a protein concentration at least 100 mg/dl.

1. Prepare (with proteins diluent solution) freshly dilution of serum sample, so that the concentration of each globulin in corresponded dilution is near 100-200 mg % . For example: when in a case the concentration of IgG globulin is 3000 mg %, then the sample should be diluted 1:16 for IgG (=187mg% ). In case of sample with globulin concentration near the normal level, or with unknown concentration, dilute the sample 1:6(1 vol. serum +5 vol. protein diluent) for PE position and 1:9 for all other positions (1 volume serum +8 volume diluent).
2. Prepare all the reagents (staining-destaining solution) according to the instructions and fill the staining and destaining tank.
3. Fill the distilled water tank with distilled water and 100 ml of tap water.
4. Check that the waste tank is empty.
5. Turn on your computer.
6. Turn on the general switch located on the left backside of the instrument.
7. Then press the power button on the front of the machine (Blue button).
8. Lift the cover of the machine.
9. Take out the sponges from their packaging. Add 6.5 ml buffer on each sponge and apply them on the migration chamber slots.
10. Locate the migration chamber in correct position.
11. Take the agarose gel out of its packaging, uncover it from the plastic plate and put it on the backside of the plate in horizontal position.
12. Blot the gel for 5" with a gel blotter strip in the zone of sample - application,
13. Place the sample template on the application zone carefully. Rub the template with forefinger gently to eliminate trapped air bubbles.
14. Using a 5-µl pipette, apply 5 µl of serum dilution across each corresponding slit [The application of the samples should be done as quickly as possible. The application slits should not be allowed to dry].
15. Let the samples absorb into the gel for 2 minutes and then blot the superfluous sample with a gel blotter strip.
16. Remove both the sample template gently and gel blotter strip and discard them.
17. Apply 0,5ml distilled water on the electrophoresis area.
18. Remove the gel holder and the gel hold support.
19. Fasten the film on the gel holder on the side marked + (anode). Apply the film on the electrophoresis area.
20. Apply carefully the gel hold support on the gel.
21. Double Click on Gemini software.
22. Click on LOGIN.
23. Click on METHODS.
24. Select the type of electrophoresis (serum).
25. Adjust the parameters:

IFEAD01				
Phase	Time	°C	mA	
Premigration	3'	20	-	
Sampling samples	20"	-	-	
Deposition samples	20"	-	-	
Migration	9'	20	190	
Desiccation	11'	75	-	
Staining	4'	-	-	
Destaining (9 cycles)	6'09"	-	-	

26. Save the parameters.
27. Click on the option New Analysis.
28. Select serum protein electrophoresis.
29. Click on the option Big Strip.
30. Select one patient and click on the option Confirm.
31. Enter the patient's data. Click on the option SAVE.
32. Click on the option START ANALYSIS.
33. Check the status of the machine.
34. Click on the option Continue the process.
35. After the migration phase (9 min) stop the process manually.  
**Warning! Allow the robotic arm to return the migration chamber to the parking position. Then click on the option Stop.**
36. Take the gel and apply it on the back side of the plastic plate
37. Blot the gel surface with a gel blotter sheet and apply on the gel surface the antiserum template.
38. Rub the template with forefinger gently to eliminate air bubbles.
39. Apply into the corresponding troughs of the antisera template:

Position	Reagents / Volume
PE	PFS / 70 µl
IgG, IgA, IgM, κ, λ	Antisera / 40 µl

40. Incubate the gel for 10 minutes.
41. Remove the antisera template and discard it.
42. Put on the gel one gel blotter sheet and one drying blotter sheet; place on the gel a development weight (about 2 kg) for 2 minutes.
43. Soak the gel in saline solution (0.9% NaCl) for 5 minutes.
44. Repeat step 42 and 43 three more times.
45. Fasten the film on the gel holder on the side marked + (anode). Apply the film on the electrophoresis area.
46. From the menu bar on the left side of the screen click on the option MACHINE STATUS.
47. Click on the option User Service. Click on the option RESUME PROCESS SPE FROM EVAPORATION.
47. The instrument will complete the drying, staining and destaining process automatically.
48. Evaluate visually the results (see conclusion / troubleshooting).

## Results:

The identification and the determination of the monoclonal band should be done by comparing the location of the band(s) with the band(s) with the same location in the PE position (see conclusion / troubleshooting).

## Expected values and Evaluation of Results:

In case of monoclonal paraproteinemia monoclonal band(s) will be appear with one or more anti heavy and /or light chain globulin. If you get monoclonal band(s) in PE position, the conclusion / troubleshooting should be as follow:

PE	Anti IgG	Anti IgA	Anti IgM	Anti κ	Anti λ	Conclusion/ Troubleshooting
	+	-	-	-	-	Monoclonal IgG heavy chain/-
	-	-	-	+	-	Monoclonal IgG κ- chain/-
	-	-	-	-	+	Monoclonal light κ- chain/ Think about IgD and IgE too.
	+	-	-	-	-	Monoclonal IgG λ-chain
	-	+	-	-	-	Monoclonal IgA κ- chain
	-	+	-	-	+	Monoclonal IgA λ- chain
	+	+	-	-	+	Biclonal IgG, IgA IgA λ- chain
	-	-	+	-	-	Monoclonal IgM heavy chain
	-	-	+	-	+	Monoclonal IgM λ- chain
	+	+	+	+	+	Fibrinogen or Heavy chain of IgD or IgE. Do not use plasma for electrophoresis! Examine the case with Anti IgD and IgE antisera too.
	+	+	+	+	+	If cryoglobulins, rheumatoid factor, or immuno complexes are present in sample monoclonal bands appear with more than one or with all antisera.

**Sensitivity:** The sensitivity of fixation solution (on PE position) is 10X higher than with Amido Black. The antisera can detect the specific antigen in a concentration of 80-400mg%.

**Specificity:** The antisera are monospecific and they do not react with fibrinogen or other human proteins.

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