The two major variant hemoglobins in the U.S. in terms of frequency and pathology are HbS and HbC. The early diagnosis of sickle cell disease (HbSS, HbSC, HbSD, and HbS/S-thal) is crucial to treat against severe infections. Homozygous HbC and the heterozygous traits HbA, HbC, HbAE, and HbAO cause mild hemolytic anemia. The diagnosis of their presence is also important for genetic counseling. The thalassemias are quantitative disorders in which the globin chain producing cells are diminished or absent. Alpha-thalassemia trait is characterized by the presence of Hb Barts (gamma 4) in newborns and HbH (Beta 4) in adults. Beta-thalassemia major is characterized by the lack of HbA and the presence of HbF and HbA2. Beta-Thalassemia minor is characterized by having elevated HbA2 and reduced levels of HbA.

Another quantitative disorder is hereditary persistent fetal hemoglobin (HPFH), in which HbF remains present in adults.

BIBLIOGRAPHY


SPIFE® Hemoglobin IEF System

Cat. No. 3428
SPIFE Hemoglobin IEF Gels (10) 3704
Hemoglobin Reagent (1 x 25 mL) 1088

Other Supplies and Equipment

The following items, needed for performance of the SPIFE Hemoglobin IEF Procedure, must be ordered individually:

- Cat. No
- SPIFE IEF Electrodes and Adapters 3704
- SPIFE 3000 Analyzer 1088
- REP Prepper 1359
- AFSC Control 5331
- AA Control 5328
- SPIFE IgG IEF Square Electrode 3703

For Sales, Technical and Order Information and Service Assistance, call 800-231-5663 toll free.

Helena Laboratories warrants its products to meet our published specifications and to be free from defects in materials and workmanship. Helena’s liability under this contract or otherwise shall be limited to replacement or refund of any amount not to exceed the purchase price attributable to the goods as to which such claim is made. These alternatives shall be buyer’s exclusive remedies.

In no case will Helena Laboratories be liable for consequential damages even if Helena has been advised of the possibility thereof.

The foregoing warranties are in lieu of all warranties expressed or implied, including, but not limited to, the implied warranties of merchantability and fitness for a particular purpose.

Interfering Factors:

1. Under very low ambient humidity, the gel may exhibit drying, resulting in skewed bands and uneven band migration. Electrophoresis at a slightly lower temperature will eliminate this problem.
2. A band will migrate in an arched manner if the concentration of hemoglobin is too high for that band. Resolution should be adequate, but the phenomena can be prevented by application of less sample. A sample application of 1 µL should rectify the problem.

PREPARATION FOR USE:

The SPIFE Hemoglobin IEF system is intended for the qualitative identification of hemoglobin variants using isoelectric focusing on the SPIFE 3000. The system is used to aid in the diagnosis of hemoglobinopathies.

SUMMARY

Hemoglobins (Hb) are composed of polypeptide chains, called globin, and iron protoporphyrin heme groups. A specific sequence of amino acids constitutes each of four polypeptide chains. Each normal hemoglobin molecule contains one pair of alpha and one pair of non-alpha chains. In normal adult hemoglobin (HbA), the non-alpha chains are called beta. The non-alpha chains of fetal hemoglobin are called gamma. A minor (3%) hemoglobin fraction called HbA2 contains alpha and delta chains. Two other chains are formed in the embryo.

The major hemoglobin in the erythrocytes of the normal adult is HbA, and there are small amounts of HbA2 and HbF. In addition, over 400 mutant hemoglobins are now known, some of which may cause serious clinical effects, especially in the homozygous state or in combination with another abnormal hemoglobin. Wintrobe’s divides the abnormalities of hemoglobin synthesis into three groups: (1) production of an abnormal protein molecule (e.g. sickle cell anemia), (2) reduction in the amount of normal protein synthesis (e.g. thalassemia), and (3) development anomalies (e.g. hereditary persistence of fetal hemoglobin (HPFH)).

The two most common hemoglobin variants commonly seen in the United States are HbS and HbC. Hemoglobinopathies are best diagnosed by other methods, such as globin chain electrophoresis described by Schneider. Pioneering work by Vesterberg and Svensson in the identification of hemoglobinopathies. Other methods of technology have been described by Drysdale, Righetti, and Bunn; Basset, Beuzard, Gare, and Rosa; Monte, Beuzard, and Rosa; and Galacteros, Klem, Caburi-Martin, Rosa, and Lubin. Because the isoelectric point of any protein is an absolute value, isoelectric focusing may reveal many more abnormal hemoglobins than conventional electrophoretic or column methods which, when combined with interpretation of clinical data, minimizes the chance of missing a “silent” variant.

PRINCIPLE

By using ampholyte buffers appropriate for isoelectric focusing of hemoglobin (pH 6-8), the separation and identification of many abnormal hemoglobins, indistinguishable from other more common forms on alkaline and acidic electrophoresis, are possible since the migration is only affected by the isoelectric point of the protein. 

Hemoglobins with a pH difference of less than 0.05 pH units could be resolved.

REAGENTS

1. SPIFE Hemoglobin IEF Gel

Ingredients: Each gel contains 1% w/v agarose, 5.3% v/v carrier ampholytes, and 0.01% thymol as a preservative.

WARNING: FOR IN-VITRO DIAGNOSTIC USE ONLY.

Preparation for Use: The gels are ready for use as packaged.

Storage and Stability: The gels should be stored horizontally at 15 to 30°C, in the protective packaging, and are stable until the expiration date indicated on the package. DO NOT FREEZE THE GELS OR EXPOSE THEM TO EXCESSIVE HEAT.

Signs of Deterioration: Discard the gel if any of the following conditions occur, indicating deterioration of the gel: (1) crystalline appearance indicating the agar has been frozen, (2) cracking and peeling indicating drying of the agar, (3) bacterial growth indicating contamination.

2. Hemolysate Reagent

Ingredients: The reagent is an aqueous solution containing 0.07% potassium cyanide, 0.005 M (0.18%) ethylenediaminetetraacetate (EDTA), and 0.175% saponin.

WARNING: FOR IN-VITRO DIAGNOSTIC USE ONLY. NEVER PIPETTE BY MOUTH. The reagent contains potassium cyanide.

Preparation for Use: The reagent is ready for use as packaged.

Storage and Stability: The reagent should be stored at 15 to 30°C and is stable until the expiration date indicated on the vial.

Signs of Deterioration: The reagent should be a clear, pale yellow solution.

INSTRUMENT

A SPIFE 3000 unit (Cat. No. 1088) is used to electrophorese and dry the gels. Refer to the Operator’s Manual for detailed instructions.
**SPECIMEN COLLECTION AND HANDLING**

**Specimen:** The specimen of choice is whole blood collected in EDTA tubes. However, washed, packed cells may also be used.

**Specimen Storage:** Whole blood specimens and packed cells may be stored at 2 to 8°C for one week.

**Specimen Preparation:** If normal hemolysates must be prepared for each patient sample:

1. **Whole Blood Samples**
   - a. Prepare the patient sample lysate using a 1:4 dilution with Hemolysate Reagent. Mix one part (25 µL) of whole blood with three parts (75 µL) Hemolysate Reagent.
   - b. Vortex briefly and allow the sample to stand at least 5 minutes prior to use. It is important that the cells be completely lysed (forming a clear red hemolysate) before applying the sample to the gel.  
   - c. Wash packed cells 3 times by resuspending in 5 to 10 volumes of normal saline solution (0.85% NaCl), centrifuging and aspirating supernatant as before.
   - d. Make a 1:8 dilution by mixing one part (10 µL) washed, packed cells to seven parts (70 µL) Hemolysate Reagent.
   - e. Vortex briefly, and allow the sample to stand at least 5 minutes prior to use. It is important that the cells be completely lysed (forming a clear red hemolysate) before applying the sample to the gel.

2. **Washed, Packed Cell Samples**
   - Prepare washed, packed cells from whole blood.
     - a. Centrifuge anticoagulated blood for 10 minutes to separate cells from plasma.
     - b. Remove plasma.
     - c. Wash packed cells 3 times by resuspending in 5 to 10 volumes of normal saline solution (0.85% NaCl), centrifuging and aspirating supernatant as before.
     - d. Make a 1:8 dilution by mixing one part (10 µL) washed, packed cells to seven parts (70 µL) Hemolysate Reagent.
     - e. Vortex briefly, and allow the sample to stand at least 5 minutes prior to use. It is important that the cells be completely lysed (forming a clear red hemolysate) before applying the sample to the gel.

3. **Controls**
   - AA (Cat. No. 5328): no dilution is necessary
   - AFSC (Cat. No. 5331): 1:2 (1 part control + 1 part Hemolysate Reagent)

**PROCEDURE**

**Materials provided:** The following materials are provided in the SPIFE Hemoglobin IEF Kit (Cat. No. 3428).

- SPIFE Hemoglobin IEF Gels (10)
- Hemolysate Reagent (1 x 25 mL)

**Materials needed but not contained in the kit:**

- Cat. No.
  - SPIFE IEF Electrodes and Adapters
  - SPIFE 3000 Analyzer
  - REP Preparer
  - AFSC Control
  - SPIFE IgG IEF Square Electrode

**MATERIALS needed but not provided:**

- 0.85% NaCl

**STEP-BY-STEP METHOD**

**NOTE:** If a SPIFE procedure requiring a stain has been run prior to running the IEF gels, the stainer unit must be cleaned/washed before washing the IEF gel.

The software has an automatic wash cycle prompted by initiation of a test which does not use the stainer unit for staining when the previous test did use the stainer for

**Pretreatment:** To verify the status, press the TEST SELECT/CONTINUE button on the stainer unit to initiate the appropriate test.

**Rinse:** Place an empty Gel Holder in the stainer unit. If cleaning is required, the “Wash 1” prompt will appear, followed by “Plate out, Holder in” prompts. Press “Continue” to begin the stainer wash. The cleaning process will complete automatically in about 7 minutes. The unit is then ready to process the gel after electrophoresis.

**Gel Preparation**

1. Carefully open one end of the pouch, remove the gel from the protective packaging, and discard the overlay.

2. Use the REP Preparer to remove excess moisture from the sample wells.

3. Do NOT use REP Prep with this procedure.

4. Place left edge of gel over the chamber, aligning the round hole on the left pin. Gently lay the gel down on the chamber, starting from the left side and ending on the right side, fitting the ovoid hole over the right pin.

5. Use a lint-free tissue to wipe around the edges of the gel.

6. Insert the SPIFE IEF Electrode Adapter marked FRONT between the two magnetic posts located at the front of the chamber floor. Insert the SPIFE Electrode Adapter marked REAR between the two magnetic posts located at the back of the chamber floor.

7. Clean and wipe three IEF electrodes with a lint-free tissue.

8. Place a round electrode into the slots created by the adapter – one on each outside edge. Place the square electrode in the middle. Be sure all three electrodes are seated firmly against the gel and electrode posts.

9. Close the chamber lid and press the TEST SELECT/CONTINUE button until an open USER TEST appears on the display.

**II. Sample Application and Electrophoresis**

1. Using the instructions provided in the SPIFE 3000 Operator’s Manual, set up the parameters as follows:
   - Due to variation in environmental conditions, a dry time of 20 minutes is recommended, but it may range from 20 to 30 minutes.

2. **Electrophoresis Unit**
   - **No Prompt**
     - **Pause 1**
       - **00:30**
       - **14°C**
   - **To Continue**
     - Electrophoresis 1
       - **4.00**
       - **14°C**
       - **650 V**
       - **50 mA**
   - **Prompt**
     - **END OF TEST**

3. **Stainer Unit**
   - **No Prompt**
     - **Dry 1**
       - **00:00**
       - **70°C**
     - **No Prompt**
       - **END OF TEST**

4. Press the START/STOP button. An option to either begin the test or skip the operation will be presented. Press START/STOP to begin electrophoresis. When completed, the SPIFE 3000 will beep.

5. Remove the chamber lid, and remove the electrodes and adapters.

6. Immediately remove the gel from the electrophoresis chamber.

7. Remove the Gel Holder from the stainer chamber. Attach the gel to the holder by placing the round hole in the gel mylar over the left pin on the holder and the ovoid hole over the right pin on the holder.

8. Place the Gel Holder with the attached gel facing backwards into the stainer chamber.

9. With the appropriate test name on the display, press the START/STOP button. An option to either begin the test or skip the operation will be presented. Press START/STOP to begin. The instrument will dry the gel.

10. When the process is completed, the instrument will beep. Remove the Gel Holder from the stainer and the gel from the holder.

**III. Evaluation of the Hemoglobin Bands**

The hemoglobin gels may be visually inspected for the presence of hemoglobin bands. The Helena Hemo Controls provide a marker for band identification.

**Stability of End Product**

Gels should be examined the same day of electrophoresis.

**Quality Control**

Use of the AFSC Control (Cat. No. 5331) and AA Control (Cat. No. 5328) is recommended with each run. Dilute the AFSC Control 1:2 with Hemolysate Reagent before use.

**Caution:** Bands from fresh whole blood may electrophorese at a slightly slower rate than the control. Older samples may electrophorese at a slightly faster rate than the control. If controls do not perform as expected, test results should be considered suspect or invalid.

**RESULTS**

Figure 1 shows the relative positions of commonly seen abnormal hemoglobins on the SPIFE Hemoglobin IEF Gel. HbE and Hb-Arab migrate slightly anodal to HbA and cannot be separated, as cannot HbN-Baltimore and Hb-Texas. Confirmation by citrate agar or alkaline acetate electrophoresis is recommended. Globin chain analysis and structural studies are recommended as a last resort. It is impossible to distinguish HbSS, HbSβ/β- thalassemia, and HbS/HbF in neonatal. The presence of HbS+HBF requires family studies. HbCC cannot be distinguished from HbCβ/β-thalassemia, and HbE cannot be distinguished from HbEβ/β-thalassemia. Both require family studies.

Beta-Thalassemia major cannot be distinguished from homozgyous HPFHV or normals with low concentrations of HSA by electrophoresis and require clinical and family studies.*

Gels which do not lay flat in the chamber or those with surface artifacts should not be used.

**Interfering Substances:**

The presence of ferric-ferrous or chloroform extraction of lipids is not recommended. Especially troublesome are ferric-ferrous hybrids, which appear as twin bands cathodal to the parent molecule. In order to use badly hemolyzed or aged specimens, they may be centrifuged, the plasma removed and discarded, and the red cells washed with physiological saline twice to minimize interfabinet bands. Washed specimens may be used as packed cells or diluted with physiological saline and treated as whole blood.

Lipemic samples may give erroneous results. Toluene or chloroform extraction of lipids is not recommended. If using lipemic samples, removal of the plasma by salting out is sufficient for satisfactory performance. Elevated levels of plasma proteins may cause water droplets to appear over the sample application area, causing diffusion of nearby hemoglobin bands. In severe cases, removal of the plasma and washing the red cells with saline will eliminate the problem.
PROCEDURE

Materials provided: The following materials are provided in the SPIFE Hemoglobin IEF Kit (Cat. No. 3428).

Individual items are not available:
- SPIFE Hemoglobin IEF Gels (10)
- Hemolysate Reagent (1 x 25 mL)

Materials needed but not contained in the kit:
- AFSC Electrodes and Adaptor
- SPIFE 3000 Analyzer
- REP Preparer
- AFSC Control
- SPIFE IgG IEF Square Electrode
- NaCl

Materials needed but not provided:
- 0.85% NaCl

STEP-BY-STEP METHOD

NOTE: If a SPIFE procedure requiring a stain has been run prior to running the IEF gels, the stainer unit must be cleaned/washed before loading the IEF gel.

The software has an automatic wash cycle prompted by initiation of a test which does not use the stainer unit for staining when the previous test did use the stainer for washing. To avoid delays after electrophoresis, this wash cycle should be initiated at least seven minutes prior to the end of the staining period. To verify the status, press the TEST SELECT/CONTINUE button on the stainer until the appropriate test is selected. Place an empty Gel Holder in the stainer unit. If cleaning is required, the “Wash 1” prompt will appear, followed by “Plate out, Holder in” prompts. Press “Continue” to begin the stainer wash. The cleaning process will automatically in about 7 minutes. The unit is then ready to process the gel after electrophoresis.

I. Gel Preparation
1. Carefully open one end of the pouch, remove the gel from protective packaging, and discard the overlay.
2. Use the REP Preparer to remove excess moisture from the sample wells.
3. Do NOT use REP Prep with this procedure.
4. Place left edge of gel over the chamber, aligning the round hole on the left. Gently lay the gel down on the chamber, starting from the left side and ending on the right side, fitting the obround hole over the right pin.
5. Use a lint-free tissue to wipe around the edges of the gel.
6. Insert the SPIFE IEF Electrode Adapter marked FRONT between the two magnetic posts located at the front of the chamber floor. Insert the IEF Electrode Adapter marked REAR between the two magnetic posts located at the back of the chamber floor.
7. Clean and wipe the three IEF electrodes with a lint-free tissue.
8. Place a round electrode into the slots created by the adapter – one on each outside edge. Place the square electrode in the middle. Be sure all three electrodes are seated firmly against the gel and electrode posts.
9. Close the chamber lid and press the TEST SELECT/CONTINUE button until an open USER TEST appears on the display.

II. Sample Application and Electrophoresis
1. Using the instructions provided in the SPIFE 3000 Operator’s Manual, set up the parameters as follows:
   - Due to variation in environmental conditions, a dry time of 20 minutes is recommended, but it may range from 20 to 30 minutes.

Electrophoresis Unit

- No Prompt
- Prompt
- Prompt
- Prompt
- Prompt
- Prompt

II. RESULTS

Figure 1 shows the relative positions of commonly seen abnormal hemoglobins on the SPIFE Hemoglobin IEF Gel. HbE and HbA2-Asian migrate slightly anodal to HbA, which itself is slightly anodal to HbC. HbG-Philadelphia and HbD-Punjab are clearly separated from HbS, as is Hb Lepore. HbA, HbF, and HbF (acyethylated HbF) are clearly separated from each other, allowing identification between heterozygous AS trait, homozygous SS anemia, and HbS/HbS thalassemia.

Anodal, or fast, hemoglobins such as Bart’s, HbH, HbJ, and HbN-Baltimore may not be separated by conventional electrophoresis but are clearly separated by isoelectric focusing.

II. LIMITATIONS

Some abnormal hemoglobins have similar isoelectric points and cannot be distinguished using isoelectric focusing. Examples indistinguishable from HbS are HbG-Galveston and HbG-Norfolk. The variants Hb Hammersmith, Brigham, and Bethesda are indistinguishable from HbA, HbE, HbC-Harlem, HbD-Arab, and Hb-Koln cannot be separated, as cannot HbN-Baltimore and HbN-Texas. Confirmation by citrate agar or alkaline acetate electrophoresis is recommended. Globin chain analysis and structural studies are recommended as a last resort. It is impossible to distinguish HbS, HbS/HbA2 thalassemia, and HbS/HbF in neonatal. The presence of HbS+HbF requires family studies. HbCC cannot be distinguished from HbC/HbF thalassemia, and HbE cannot be distinguished from HbE/HbA2 thalassemia. Both require family studies. Beta-Thalassemia major cannot be distinguished from homozgyous HbF/HbF or normals with low concentrations of HbA by electrophoresis and require clinical and family studies.

Gels which do not lay flat in the chamber or those with surface artifacts should not be used.

Interfering Substances:

The use of fresh whole blood is recommended to avoid artifactual bands caused by oxidation or denaturation of the hemoglobin. Especially troublesome are ferri-ferrois hybrids, which appear as twin bands cathodal to the parent molecule. In order to use badly hemolyzed or aged specimens, they may be centrifuged, the plasma removed and discarded, and the red cells washed with physiologically saline twice to minimize artifactual bands. Washed specimens may be used as packed cells or diluted with physiologically saline and treated as fresh whole blood.

Lipemic samples may give erroneous results. Toluene or chloroform extraction of lipids is not recommended. If using lipemic samples, removal of the plasma by salting out is sufficient for satisfactory performance. Elevated levels of plasma proteins may cause water drops to appear on the sample application area, causing difficulty of seeing nearby hemoglobin bands. In severe cases, removal of the plasma and washing the red cells with saline will eliminate the problem.
Interfering Factors:
1. Under very low ambient humidity, the gel may exhibit drying, resulting in skewed bands and uneven band migration. Electrophoresis at a slightly lower temperature will eliminate this problem.
2. A band may migrate in an arched manner if the concentration of hemoglobin is too high for that band. Resolution should be adequate, but the phenomena can be prevented by application of less sample. A sample application of 1 µL should rectify the problem.

PREPARATION OF THE GELS

The gels should be stored at 4°C. Each gel contains 1% w/v agarose, 5.3% NaCl, and 0.01% thymol. The gels are ready for use as described in the Hemoglobin IEF Procedure. The reagent should be a sterile glass suspension of agar. If the agar contains free air bubbles, these should be removed, and the gel acted upon Immunohematologica by application of less sample. A sample application of 1 µL should rectify the problem.

PREPARATION FOR USE

The following items, needed for performance of the SPIFE Hemoglobin IEF Procedure, must be ordered individually.

**Cat. No.**
1. SPIFE® Hemoglobin IEF Gels (10) 3428
2. Hemoglobin IEF Reagents (1 x 25 mL) 3503
3. Other Supplies and Equipment 3704

For Sales, Technical and Order Information and Service Assistance, call 800-231-5663 toll free.

**BIBLIOGRAPHY**


**SUMMARY**

Hemoglobin (Hb) is composed of polypeptide chains, called globin, and iron protoporphyrin heme groups. A specific sequence of amino acids constitutes each of four polypeptide chains. Each normal hemoglobin molecule contains one pair of alpha and one pair of non-alpha chains. In normal adult hemoglobin (HbA), the non-alpha chains are called beta. The non-alpha chains of fetal hemoglobin are called gamma. A minor (3%) hemoglobin fraction called HbAβ contains alpha and delta chains. Two other chains are formed in the embryo.

The major hemoglobin in the erythrocytes of the normal adult is HbA, and there are small amounts of HbA2 and HbF. In addition, over 400 mutant hemoglobins are now known, some of which may cause serious clinical effects, especially in the homozygous state or in combination with another abnormal hemoglobin. Wintrobe® divides the abnormalities of hemoglobin synthesis into three groups: (1) production of an abnormal protein molecule (e.g. sickle cell anemia), (2) reduction in the amount of normal protein synthesis (e.g. thalassemia), and (3) development anomalies (e.g. hereditary persistence of fetal hemoglobin (HbF)).

The two major variant hemoglobins most commonly seen in the United States are HbS and HbC. Hb Lepore, HbE, HbG-Philadelphia, HbD-Los Angeles, and HbO-Arab may be seen less frequently.

Electrophoresis is generally considered the best method for screening and diagnosis of hemoglobinopathies. Routine testing of adults and newborns consists of alkaline electrophoresis followed by agar electrophoresis in order to confirm the presence of HbS or Hb-like variants (as HbG-Philadelphia and D-Los Angeles) or the presence of HbA2-HbC and HbE, which are indistinguishable from HbA on alkaline electrophoresis. The resolving power of these systems is rather poor for many abnormal hemoglobins, however, and positive identification must be determined by other methods, such as globin chain electrophoresis described by Schneider.

Pioneering work by Vesterberg and Svensson in the manufacture and use of ampholyte buffers has enabled isoelectric focusing to become a powerful diagnostic tool in the identification of hemoglobinopathies. Other methods of technology have been described by Drysdale, Righetti, and Bunn, Basset, Beuzard, Garel, and Rosa, and Pioneering work by Vesterberg and Svensson in the manufacture and use of ampholyte buffers has enabled isoelectric focusing to become a powerful diagnostic tool in the identification of hemoglobinopathies. Other methods of technology have been described by Drysdale, Righetti, and Bunn, Basset, Beuzard, Garel, and Rosa.

Because the isoelectric point of any protein is an absolute value, isoelectric focusing may reveal many more abnormal hemoglobin variants than conventional electrophoretic or column methods which, when combined with interpretation of clinical data, minimizes the chance of missing a "silent" variant.

**PRINCIPLE**

By using ampholyte buffers appropriate for isoelectric focusing of hemoglobin (pH 6-8), the separation and identification of many abnormal hemoglobins, indistinguishable from other more common forms on alkaline and acidic electrophoresis, are possible since the migration is only affected by the isoelectric point of the protein. Hemoglobins with a pH difference of less than 0.05 pH units could be resolved.

**REAGENTS**

1. SPIFE® Hemoglobin IEF Gel

**Ingredients:** Each gel contains 1% w/v agarose, 5.3% NaCl carrier ampholytes, and 0.01% thymol as a preservative.

**WARNING:** IN-VITRO DIAGNOSTIC USE ONLY. Preparation for Use: The gels are ready for use as packaged.

**Storage and Stability:** The gels should be stored horizontally at 15 to 30°C, in the protective packaging, and are stable until the expiration date indicated on the package. DO NOT FREEZE THE GELS OR EXPOSE THEM TO EXCESSIVE HEAT.

**Signs of Deterioration:** Discard the gel if any of the following conditions occur, indicating deterioration of the gel: (1) crystalline appearance indicating the agar has been frozen, (2) cracking and peeling indicating drying of the agar, (3) bacterial growth indicating contamination.

2. Hemolysate Reagent

**Ingredients:** The reagent is an aqueous solution containing 0.07% potassium cyanide, 0.005 M (0.18%) ethylenediaminetetraacetate (EDTA), and 0.175% saponin.

**WARNING:** IN-VITRO DIAGNOSTIC USE ONLY. NEVER PIPETTE BY MOUTH. The reagent contains potassium cyanide.

**Preparation for Use:** The reagent is ready for use as packaged.

**Storage and Stability:** The reagent should be stored at 15 to 30°C and is stable until the expiration date indicated on the vial.

**Signs of Deterioration:** The reagent should be a clear, pale yellow solution.

**INSTRUMENT**

A SPIFE® 3000 unit (Cat. No. 1088) is used to electrophorese and dry the gels. Refer to the Operator’s Manual for detailed instructions.