*PLEASE READ!!*

**HELENA LABORATORIES**

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We also provide the procedure in an Adobe Acrobat PDF format for download at www.helena.com as a “MASTER” file copy. Below are the specifications and requirements for using these digital files. Following the specifications is the procedure major heading sequence as given in the FDA style. Where there is a difference in order, or other notation in the outline, this will be indicated in braces { }.

WHAT YOU NEED TO KNOW:

1) These files represent the most current revision level to date. Your current product inventory could contain a previous revision level of this procedure.

2) The Microsoft Word document provides the text only from the master procedure, in a single-column format.

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4) By downloading this procedure, your institution is assuming responsibility for modification and usage.

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HELENA LABORATORIES LABELING – Style/Format Outline

1. PRODUCT {Test} NAME
2. INTENDED USE and TEST TYPE (qualitative or qualitative)
3. SUMMARY AND EXPLANATION
4. PRINCIPLES OF THE PROCEDURE

{*NCCLS lists SAMPLE COLLECTION/HANDLING next}*

1. REAGENTS (name/concentration; warnings/precautions; preparation; storage; environment; Purification/treatment; indications of instability)
2. INSTRUMENTS required – Refer to Operator Manual (... for equipment for; use or function; Installation; Principles of operation; performance; Operating Instructions; Calibration\* {\*is next in order for NCCLS – also listed in “PROCEDURE”}’ precautions/limitations/hazards; Service and maintenance information
3. SAMPLE COLLECTION/HANDLING
4. PROCEDURE

{*NCCLS lists QUALITY CONTROL (QC) next}*

9) RESULTS (calculations, as applicable; etc.)

10) LIMITATIONS/NOTES/INTERFERENCES

11) EXPECTED VALUES

12) PERFORMANCE CHARACTERISTCS

13) BIBLIOGRAPHY (of pertinent references)

14) NAME AND PLACE OF BUSINESS OF MANUFACTURER

15) DATE OF ISSUANCE OF LABELING (instructions)

For Sales, Technical and Order Information, and Service Assistance,   
call Helena Laboratories toll free at 1-800-231-5663.

Form 364

Helena Laboratories

1/2006 (Rev 3)

**QuickGel® Split Beta SPE Procedure**

**Cat. No. 3350, 3550, 3550T**

**INTENDED USE**

The QuickGel Split Beta SPE System is intended for the separation of serum, cerebrospinal fluid (CSF), or urine proteins by agarose gel electrophoresis using the SPIFE 3000 system or the QuickGel Chamber.

Cat. No. 3350 - for use with SPIFE 3000

Cat. No. 3550, 3550T - for use with QuickGel Chamber

For In Vitro Diagnostic use.

Rx Only

**SUMMARY**

Serum contains over one hundred individual proteins, each with a specific set of functions and subject to specific variation in concentration under different pathologic conditions.1 Since the introduction of moving-boundary electrophoresis by Tiselius2 and the subsequent use of zone electrophoresis, serum proteins have been fractionated on the basis of their electrical charge at a particular pH into five classical fractions: albumin, alpha1, alpha2, beta and gamma proteins. Each of these classical electrophoretic zones, with the exception of albumin, normally contains two or more components. The relative proportions of these fractions have proven to be useful aids in the diagnosis and prognosis of certain disease states.3-5

**PRINCIPLE**

Proteins are large molecules composed of covalently linked amino acids. Depending on electron distributions resulting from covalent or ionic bonding of structural subgroups, proteins can be either polar or nonpolar at a given pH. In the QuickGel Serum Protein procedures, proteins are separated according to their respective electrical charges on agarose gel using both the electrophoretic and electroendosmotic forces present in the system. The proteins are then stained with a visible stain.

**REAGENT**

**1. QuickGel Split Beta SPE Gel**  **Ingredients:** Each gel contains agarose in a tris-barbital/MOPS buffer with calcium lactate, a stabilizer, and a preservative.   
**WARNING: FOR IN-VITRO DIAGNOSTIC USE ONLY.** The gel contains barbital which, in sufficient quantity, can be toxic.

**Preparation for Use:** The gels are ready for use as packaged. **Storage and Stability:** The gels should be stored at room temperature (15 to 30°C) and are stable until the expiration date indicated on the package. The gels must be stored horizontally in the protective packaging in which they are shipped. **DO NOT REFRIGERATE OR FREEZE THE GELS.**   
**Signs of Deterioration:** Any of the following conditions may indicate deterioration of the gel: (1) crystalline appearance indicating the agarose has been frozen, (2) cracking and peeling indicating drying of the agarose, (3) bacterial growth indicating contamination, (4) thinning of the gel blocks.

**2. Acid Blue Stain   
Ingredients:** When dissolved as directed, the stain contains 0.5% (w/v) acid blue stain.   
**WARNING: FOR IN-VITRO DIAGNOSTIC USE ONLY. DO NOT INGEST.**   
**Preparation for Use:** Dissolve the dry stain (entire contents of vial) in 1 L of 5% acetic acid. Mix thoroughly for 30 minutes.   
**Storage and Stability:** The dry stain should be stored at 15 to 30°C and is stable until the expiration date indicated on the package. The diluted stain is stable six months when stored at 15 to 30°C.   
**Signs of Deterioration:** The diluted stain should be a homogeneous mixture free of precipitate. Discard if precipitate forms.

**3. Citric Acid Destain**  **Ingredients:** After dissolution, the destain contains 0.3% (w/v) citric acid. **WARNING: FOR IN-VITRO DIAGNOSTIC USE. DO NOT INGEST - IRRITANT.   
Preparation for Use:** Pour 11 L of deionized water into the Destain vat. Add the entire package of Destain. Mix well until completely dissolved. **Storage and Stability:** Store the Destain at 15 to 30°C. It is stable until the expiration date on the package.

**Signs of Deterioration:** Discard if solution becomes cloudy.

**4. Acid Violet Stain (Optional Urine Stain)**  **Ingredients:** The stain is comprised of Acid Violet stain. **WARNING: FOR IN-VITRO DIAGNOSTIC USE ONLY. DO NOT INGEST.**

**Preparation for Use:** Dissolve the dry stain in 1 liter of 10% acetic acid and mix thoroughly. Fill the SPIFE stain vat. **Storage and Stability:** The dry stain should be stored at 15 to 30°C and is stable until the expiration date indicated on the package. The stain solution is stable six months when stored at 15 to 30°C in a closed   
container.   
**Signs of Deterioration:** The diluted stain should be a homogeneous mixture free of precipitate.

**INSTRUMENT**

A SPIFE 3000 or the QuickGel Chamber must be used to electrophorese, stain, destain, and then dry the gels. The gels may be scanned on the QuickScan Touch Plus (Cat. No. 1640) or a separate densitometer. Refer to the Operator’s Manual for detailed instructions.

**SPECIMEN COLLECTION AND HANDLING**

**Specimen:** Fresh serum, CSF or urine is the specimen of choice. Use of plasma will cause a fibrinogen band to appear as a distinct narrow band between the beta and gamma fractions.

**Storage and Stability:** If storage of serum is necessary, samples may be stored covered at 15 to 30°C for 4 days, 2 to 8°C for 2 weeks, or –20°C for 6 months.6 Urine or CSF samples may be stored covered at 2 to 8°C for up to 72 hours or at –20°C for 1 month.

**Urine Sample Preparation:** Urine samples may be run diluted, neat or concentrated. Shake samples to homogenize. Centrifuge desired volume at 2000 x g for 5 minutes. Remove supernatant and concentrate as follows:

Total Protein (mg/dL) Conc. Factor

<50 100x

50-100 50x

100-300 25x

300-600 10x

>600 5x

**CSF Sample Preparation:** CSF samples may be used after proper concentration (10-50X).

**Interfering Factors:**

1. Hemolysis may cause false elevation in the alpha2 and beta fractions.

2. Inaccurate results may be obtained on specimens left uncovered, due to evaporation.

**PROCEDURE FOR SPIFE 3000**

**Materials provided:** The following materials needed for the procedure are contained in the QuickGel Split Beta SPE Kit (Cat. No. 3350). Individual items are not available.

QuickGel Split Beta SPE Gels (10)

Acid Blue Stain (1 vial)

QuickGel Blotter C (10)

Citric Acid Destain (1 pkg)

Modified Applicator Blade Assembly (10)

**Material provided but not contained in the kit:**

**ITEM CAT. NO.**

SPIFE 3000 Analyzer 1088

QuickScan Touch Plus 1640

SPIFE Modified Applicator Blades 3451

(for 20 sample application)

Applicator Blade Weights 3387

SPIFE Dispo Sample Cups (deep well) 3360

Gel Block Remover 1115

SPE Normal Control 3424

SPE Abnormal Control 3425

REP Prep 3100

SPIFE Applicator Blades (for Urine & CSF) 3450

Disposable Sample Cups (for Urine and CSF) 3369

QuickGel Dispo Cup Tray 3353

SPIFE QuickGel Electrode 1111

SPIFE QuickGel Holder 3358

SPIFE QuickGel Chamber Alignment Guide 86541003

QuickGel Accessory Kit 3426

Acid Violet Stain 552351

**Materials needed but not provided:**  
 5% acetic acid

0.85% saline

**STEP-BY-STEP METHOD**

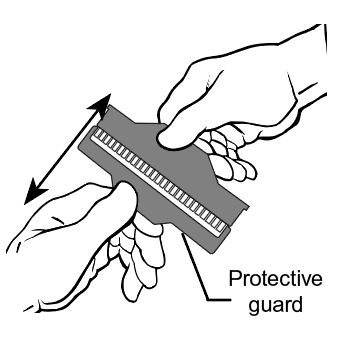
**I. Chamber Preparation**

1. The SPIFE QuickGel Chamber Alignment Guide must be used to mark the location for gel placement. It is recommended that the markings be placed directly on the copper floor under the contact sheet.

2. Remove the contact sheet and clean the chamber floor according to instructions in the Operator’s Manual.

3. Place the round hole in the guide over the left chamber pin and the obround hole over the right pin.

4. Using an indelible marker, outline the square open area onto the copper floor. Allow marking to dry, and apply another contact sheet.

**II. Sample Blade Application Method**

1. Remove one Disposable Applicator Blade from the packaging. If testing more than 10 samples, remove two Applicator Blades from the packaging. Remove the protective guard from the blades by gently bending the protective piece back and forth until it breaks free.

2. Place the Applicator Blade into the vertical slot numbered 6 in the Applicator Assembly. If using two Applicator Blades, place them into the vertical slots numbered 6 and 12. When testing serum with urine or CSF samples, serum application is made after the second urine or CSF application. Therefore the blade for serum application is not added until after the second urine/CSF application.

**NOTE: The blade assembly will only fit into the** **slots in the Applicator Assembly one way; do not try to force** **the Applicator Blades into the slots.**

3. Place an Applicator Blade Weight on top of each blade assembly.

4. Slide the Disposable Sample Cups into the top row numbered 1 to 10 of the appropriate cup tray. If testing more than 10 samples, place cups into both rows.

5. Pipette the following amount of sample into cups 1 to 5 and 6 to 10. If testing more than 10 samples, pipette sample into cups 11 to 15 and 16 to 20. Cover the tray until ready to use. **NOTE: Application of Urine and CSF samples cannot be done with the Applicator Blades or Cups packaged in the kit. Another Blade (Cat. No. 3450) and Cup (Cat. No. 3369) must be purchased.**

Sample Volume Blades Cups

Serum or control 45 mL 3451 3360

Urine/Concentrated CSF 20 mL 3450 3369

6. Carefully cut open the end of the gel pouch. Remove one gel from the protective packaging. Fold and tape the open end of the pouch to prevent drying of the gel. Remove the gel from the plastic mold and discard the mold.

7. Using a QuickGel Blotter C, gently blot the entire gel using slight fingertip pressure on the blotter. Remove the blotter.

8. Dispense approximately 1 mL of REP Prep onto the left side of the electrophoresis chamber.

9. Place the gel over the REP Prep inside the rectangle on the chamber floor. Gently lay the gel down on the REP Prep, starting from the left side and ending on the right side. Use lint-free tissue to wipe around the edges of the plastic gel backing, especially next to electrode posts, to remove excess REP Prep. Make sure the gel remains in place and that no bubbles remain under the gel.

10. Clean the QuickGel Electrodes with deionized water before and after each use. Wipe with a lint-free tissue.

11. Place a QuickGel Electrode on the outside ledge of each gel block inside the magnetic posts. Close the chamber lid and proceed to Step IV.

**III. Sample Template Application Method**

**Template application may be used for testing CSF or urine specimens which have insufficient volumes for blade application.**

1. Carefully open one end of the pouch and remove one gel from the protective packaging. Reseal the pouch with tape to prevent drying of the gel. Remove the gel from the plastic mold and discard the mold.

2. Using a QuickGel Blotter C, gently blot the entire gel using slight fingertip pressure on the blotter then remove the blotter.

3. Dispense about 1 mL of REP Prep onto the left side of the marked area of the chamber.

4. Place the gel over the REP Prep inside the rectangle on the chamber floor. Gently lay the gel down on the REP Prep, starting from the left side and ending on the right side. Use lint-free tissue to wipe around the edges of the gel backing to remove excess REP Prep. Make sure the gel remains in place and that no bubbles remain under the gel.

5. Depending on the number of samples tested, place one or two templates across the gel aligning the slits with the arrows on the gel backing.

6. Apply fingertip pressure to each template, making sure there are no bubbles under it. **NOTE:** **If wearing rubber gloves to perform this step, place a QuickGel Blotter A over the template and then apply fingertip pressure to the template. Remove the blotter. Powder from the gloves can produce gel artifacts.**

7. Clean the electrodes with deionized water before and after each use. Wipe with a lint-free tissue.

8. Place a carbon electrode on the outside ledge of each gel block inside the magnetic posts. Close the chamber lid. Proceed to Step V.

**IV. Electrophoresis with Blade Application**   
Using the instructions provided in the appropriate Operator’s Manual, set up the parameters as follows for theSPIFE 3000. If testing only serum samples, follow the instructions marked **“**• **Serum”**. If testing serum with urine or CSF, follow instructions marked **“**• **Serum and CSF or Urine”**. The blade used for serum application will be added after the second application of urine or CSF.

**NOTE: A “Dry 1” time of 10 minutes is recommended. However, due to variations in environmental conditions, the following ranges are acceptable.**

**\*Dry 1 = 10 to 15 minutes.**

**•** **Serum**

**Electrophoresis Unit**

1) No Prompt  
 Load Sample 1 00:30 21°C SPD1

2) No Prompt  
 Apply Sample 1 1:00 21°C SPD1 LOC1

3) No Prompt  
 Electrophoresis 1 8:00 21°C 350V 60 mA

4) Remove gel blocks, (continue)  
 Dry 1 \*10:00 54°C

5) No prompt  
 END OF TEST

**•** **Serum and CSF or Urine**

**Electrophoresis Unit**

1) No Prompt  
 Load Sample 1 00:30 21°C SPD1

2) No Prompt  
 Apply Sample 1 00:30 21°C SPD1 LOC1

3) No Prompt  
 Load Sample 2 00:30 21°C SPD1

4) No Prompt  
 Apply Sample 2 00:30 21°C SPD1 LOC1

5) To Continue, (continue)  
 Load Sample 3 00:30 21°C SPD1

6) No Prompt  
 Apply Sample 3 1:00 21°C SPD1 LOC1

7) No Prompt  
 Absorb 1 1:00 21°C

8) No Prompt  
 Electrophoresis 1 8:00 21°C 350V 60 mA

9) Remove gel blocks, (continue)  
 Dry 1 \*10:00 54°C

10)   
 No Prompt  
 END OF TEST

**•** **Serum and CSF or Urine**

**Stainer Unit**

1) No Prompt 4:00 REC = OFF VALVE = 3   
 Stain 1

2) No Prompt 1:00 REC = ON VALVE = 2   
 Destain 1

3) No Prompt 1:00 REC = ON VALVE = 2   
 Destain 2

4) No Prompt 1:00 REC = ON VALVE = 2   
 Destain 3

5) No Prompt \*10:00 63°C   
 Dry 1

6) No Prompt  
 END OF TEST

1. Press the **TEST SELECT/CONTINUE** button located on the electrophoresis side of the instrument until the **QG-Serum/URINE Protein** option appears on the display. Open the chamber lid.

2. Place the Cup Tray with samples on the SPIFE 3000. Align the holes in the tray with the pins on the instrument. Close the chamber lid.

3. With **QG-Serum/URINE Protein** 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** again to begin. The SPIFE 3000 will apply the samples, ectrophorese and beep when completed. Proceed to Step VI.

4. If testing serum and CSF or urine, open the chamber lid and place the Modified Blade in place for serum application. Press **TEST SELECT/CONTINUE.** SPIFE will beep when electrophoresis is complete. Proceed to Step VI.

**V. Electrophoresis with Template Application**   
Using the instructions provided in the appropriate Operator’s Manual, set up the electrophoresis parameters as follows for the SPIFE 3000.   
**NOTE: A “Dry 1” time and an “Absorb 1” time are recommended below. However, due to variations in environmental conditions, the following ranges are acceptable.**

**\*Dry 1 = 10 to 15 minutes**

**\*\*Absorb 1 = 7 to 10 minutes**

**• Template**

1) Apply Sample to Template, (continue)  
 Absorb 1 \*\*10:00 21°C

2) Blot and Remove Template, (continue)  
 Electrophoresis 1 8:00 21°C 350V 60 mA

3) Remove Gel Blocks, (continue)  
 Dry 1 \*10:00 54°C

4) No Prompt  
 END OF TEST

1. Press the **TEST SELECT/CONTINUE** button located on the electrophoresis chamber side of the instrument until the appropriate serum protein option appears on the display. Press **START/STOP** twice.

2. Open the chamber lid. Apply urine and/or CSF by placing 3 µL of each sample onto one of the ten available slits on the Sample Template.

3. Close the chamber lid, and press the **TEST SELECT/CONTINUE** button for the electrophoresis chamber. Sample application will be timed for \*\* 10 minutes.

4. Open the chamber lid and gently blot the template with a QuickGel Blotter A and carefully remove the blotter and template. Dispose of templates as biohazardous waste.

5. Close the chamber lid, and press the **TEST SELECT/ CONTINUE** button to start electrophoresis. SPIFE will beep when electrophoresis is complete.

**VI. Visualization**

1. After electrophoresis is complete, open the chamber lid and use the Gel Block Remover to remove the gel blocks. Dispose of blades and cups as biohazardous waste. Replace the electrodes on each end of the gel to prevent curling during drying.

2. Close the chamber lid and press the **TEST SELECT/CONTINUE** button to dry the gel.

3. After the gel has been dried, open the chamber lid and carefully remove the gel from the electrophoresis chamber.

4. Remove the SPIFE QuickGel Holder from the stainer chamber. While holding the gel agarose side down, slide one side of the gel backing under one of the metal bars. Bend the gel backing so that the gel is bowed, and slip the other side under the other metal bar. The two small notches in the backing must fit over the small pins to secure the gel to the holder.

5. Place the SPIFE QuickGel Holder with the attached gel facing backwards into the stainer chamber.

6. With appropriate test nameon 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 stain, destain, and dry the gel.

7. When the process is completed, the instrument will beep. Carefully remove the SPIFE QuickGel Holder from the stainer because the metal piece on the holder will be hot. Scan the bands in a densitometer.

**PROCEDURE FOR QuickGel® CHAMBER**

The following instructions are for using the QuickGel Chamber (Cat. No. 1284) for electrophoresis.

**Materials Provided:** The following materials needed for the procedure are contained in the QuickGel Split Beta SPE Kit (Cat. No. 3550 or 3550T). Individual items are not available.

**Cat. No. 3550 Cat. No. 3550T**

QuickGel Split Beta SPE Gels (10) QuickGel Split Beta SPE Gels (10)

Acid Blue Stain (1 vial) Acid Blue Stain (1 vial)

QuickGel Blotter C (10) QuickGel Blotter C (10)

Citric Acid Destain (1 pkg) Citric Acid Destain (1 pkg)

QuickGel Modified Applicator Blades (10) QuickGel Sample Templates (10)

QuickGel Dispo Sample Cups (10) QuickGel Blotter A (10)

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

**ITEM CAT. NO.**

QuickGel Chamber 1284

QuickGel Applicator 1265

QuickGel Applicator Base 1266

QuickGel Applicator Weights 1267

QuickGel Dispo Cup Tray 1268

QuickGel Dispo Sample Cups (Deep well) 1259

QuickGel Dispo Sample Cups (Shallow well) 1269

QuickGel Modified Applicator Blades 1271

QuickGel Applicator Blades (for urine and CSF) 1270

QuickGel Applicator Kit (includes

Applicator, Applicator Base, Weights, and Cup Tray) 1274

QuickGel Accessory Kit 3426

QuickGel Gel Block Remover 1262

QuickScan Touch Plus 1640

REP Prep 3100

SPE Normal Control 3424

SPE Abnormal Control 3425

Acid Violet Stain 552351

**Materials needed but not provided:**

5% acetic acid

0.85% saline

Power Supply capable of providing at least 400 Volts.

**STEP BY STEP METHOD**

**NOTE:** **The use of templates for sample application is offered as an option instead of the Applicator. Instructions are provided for both methods in Section II.**

**I. Chamber Preparation**

1. The QuickGel Chamber must be plugged into a power supply. Set a timer for 8:00 minutes and the power at 400 Volts. \*An electrophoresis time of 7:30 to 8:30 minutes is acceptable.

2. Snap the Electrophoresis Lid into place on the chamber.

3. Ensure that the chamber floor is cool (room temperature) before starting the test.

**II. Sample Application**

**A. QuickGel Applicator**

**NOTE: Application of Urine and CSF samples cannot be done with the Applicator Blades or Cups packaged in the kit. Another Blade (Cat. No. 1270) and Cup (Cat. No. 1269) must be purchased.**

Sample Volume Blades Cups

Serum or control 45 μL 1271 1259

Urine/ConcentratedCSF 20 μL 1270 1269   
Specimens with insufficient volumes may be run using the QuickGel Accessory Kit (Cat. No. 3426).

1. Remove one QuickGel Applicator Blade from the packaging. If testing more than 10 samples, remove two Applicator Blades from the packaging.

2. Urine and CSF samples must be applied on the gel three times. When testing serum with urine or CSF samples, serum application is made after the second urine or CSF application. Therefore the blade for serum application is not added until after the second urine/CSF application.

3. Place an Applicator Weight on top of each Applicator Blade. If using two Applicator Blades, place them into the vertical slots A and C of the Applicator. One blade should be placed into the slot corresponding to cup placement.

4. Place the appropriate number of QuickGel Disposable Sample Cups into either or both Row A and Row C of the Dispo Cup Tray. Pipette the appropriate amount of specimen into the Sample Cups and cover tray until ready to use. When ready, place the Dispo Cup Tray into the Applicator Base.

5. Carefully cut open one end of the gel pouch. Remove one gel from the protective packaging. Fold and tape the open end of the pouch to prevent drying of the remaining gel. Remove the gel to be used from the plastic mold and discard the mold.

6. Dispense approximately 1 mL of REP Prep onto the left side of the electrophoresis chamber.

7. Place the left notch of the gel so that it fits the left pin of the chamber floor and gently roll the gel to the right side fitting the right notch to the right pin of the chamber floor. Use a lint free tissue to wipe around the edges of the gel backing to remove any excess REP Prep. Make sure that no bubbles remain under the gel.

8. Using a QuickGel Blotter C, gently blot the entire gel using slight fingertip pressure on the blotter. Remove the blotter.

9. If testing only serum samples, follow steps 10 through 15. If testing serum and urine or CSF on the same gel, perform steps 10 through 14 twice for the urine and CSF samples. Place the Modified Blade for serum samples into the applicator, and repeat steps 10 through 14 for a total of 3 applications for urine and CSF samples and 1 application for serum samples.

10. While holding the white Applicator knob up, place the Applicator into the designated slot on the Applicator Base aligning the small red dots on the Applicator with those on the Base.

11. Slowly lower the Applicator Knob allowing the blades to enter the sample cups, and immediately start a timer for 30 seconds.

12. After 30 seconds, lift the Applicator Knob. Immediately place the Applicator into the slot on the chamber floor, aligning the red dots.

13. Slowly lower the Applicator Knob to apply sample to the gel. Set a timer for 30 seconds.

14. After the 30 seconds application, raise the Applicator Knob and remove the Applicator from the chamber.

15. Close the lid, press the power switch to turn on the chamber and start the power supply. Proceed to Step III.

**B. Sample Template Application**

1. Serum specimens and controls are diluted 1:4 (1 part serum with 3 parts 0.85% saline). Urine samples may be run diluted, neat or concentrated. CSF samples must be concentrated as instructed in **specimen collection and handling.**

2. Carefully cut open one end of the gel pouch. Remove one gel from the protective packaging. Fold and tape the open end of the pouch to prevent drying of the remaining gel. Remove the gel to be used from the plastic mold and discard the mold.

3. Dispense approximately 1 mL of REP Prep onto the left side of the electrophoresis chamber.

4. Place the left notch of the gel so that it fits the left pin of the chamber floor and gently roll the gel to the right side fitting the right notch to the right pin of the chamber floor. Use a lint free tissue to wipe around the edges of the gel backing to remove any excess REP Prep. Make sure that no bubbles remain under the gel.

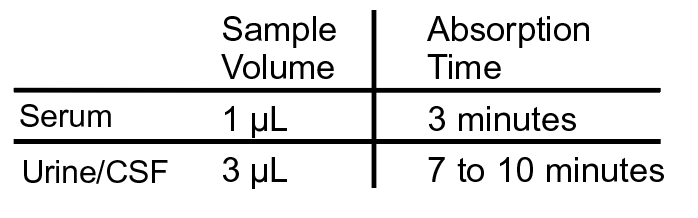
5. Using a QuickGel Blotter C, gently blot the entire gel using slight fingertip pressure on the blotter. Remove the blotter.

6. Remove one QuickGel Sample Template from the package if only one row of samples is tested. Remove two templates if two rows are tested. Hold the template so that the small hole in the corner is toward the front right side of the chamber.

7. Carefully place the template on the gel aligning the template slits with the marks on each side of the gel backing. The center hole in the template should align with the indention in the center of the gel.

8. Apply slight fingertip pressure to the template making sure there are no bubbles under it. **NOTE: If wearing rubber gloves to perform this step, place a QuickGel Blotter A over the template and then apply fingertip pressure to the template. Remove the blotter. Powder from the gloves can produce gel artifacts.**

9. Use the following chart to apply the appropriate serum dilution or urine/CSF concentration to the template slits. After the last sample application, allow time for the proper absorption. If two rows of samples are tested, start the timer for blotting the first row before applying samples to the second row. Then time the blot of the second row.



10. Use the QuickGel Blotter A to gently blot the excess sample from the template. Carefully remove the blotter and the template. Dispose of templates as biohazardous waste.

11. Close the lid, press the power switch to turn on the chamber and start the power supply.

**III. Electrophoresis and Staining**

1. Electrophorese the gel for \*8:00 minutes at 400 Volts.

2. Turn off the Power Supply and the QuickGel Chamber.

3. Using the Gel Block Remover, remove the two gel blocks from the gel. Use a lint free tissue to wipe around the edges of the gel backing to remove any excess moisture.

4. Replace the Electrophoresis Lid with the Drying Lid. Clean the two electrodes on the Electrophoresis Lid with deionized water after each use. Wipe with a lint-free tissue. Close the lid.

5. Turn on only the QuickGel Chamber. Dry the gel for 15 minutes or until dry. When drying is complete, turn the chamber off and remove the gel.

6. Fill a container with prepared stain. Fill another container with Destain solution.

7. Place the gel into the Staining Dish containing the prepared stain for 4 minutes. Remove the gel from the stain and allow it to drain on a blotter.

8. Destain the gel in three consecutive washes of Destain solution. Use a gentle alternately rocking and swirling technique. Allow the gel to remain in each wash for 1 minute. The gel background should be completely clear. Tap the gel to remove the excess destain solution.

9. Ensure that the chamber floor is clean. Replace the gel onto the QuickGel Chamber floor. Close the Drying Lid and turn on the QuickGel Chamber. Dry the gel for 10 minutes or until dry. Turn off QuickGel Chamber and remove the gel.

**Evaluation of the Protein Bands**

**Qualitative evaluation:** The urine and CSF samples run on the QuickGel Split Beta SPE Gel can only be visually inspected for the presence of the bands.

**Quantitative evaluation:** Scan the gel agarose side up. A slit size of 5 is recommended. If a QuickScan Touch Plus is used, scan on the acid blue setting.

**Stability of End Product:** The completed, dried QuickGel Split Beta SPE Gel is stable for an indefinite period of time.

**Quality Control**

SPE Normal Control (Cat. No. 3424) and SPE Abnormal Control (Cat. No. 3425) may be used to verify all phases of the procedure and should be used on each gel run. If desired, a control or patient sample may be diluted 1:7 with 0.85% saline (1 part sample + 6 parts saline) and run with CSFs for qualitative comparison. Refer to the package insert provided with the control for assay values.

**REFERENCE VALUES**

The reference range presented was established with the QuickGel Split Beta SPE System on 50 normal specimens using the SPIFE 3000 Analyzer. Each laboratory should perform its own normal range study. These values are presented as a guideline.

% of Total Protein

Protein Fraction X ± 2 S.D.

Albumin 47.6 - 61.9

Alpha1 1.4 - 4.6

Alpha2 7.3 - 13.9

Beta 10.9 - 19.1

Gamma 9.5 - 24.8

**Variations of Expected Values5**

Studies show that values are the same for both males and nonpregnant females. (Some differences are seen in pregnant females at term and in women on oral contraceptives.) Age has some effect on normal levels. Cord blood has decreased total protein, albumin, alpha2, and beta fractions, slightly increased alpha1 and normal or increased gamma fractions (largely of maternal origin). The gamma globulins drop rapidly until about three months of age, while the other fractions have reached adult levels by this time. Adult levels of the gamma globulins are not reached until 16 years of age. The albumin decreases and beta increases after the age of 40.

**RESULTS**

Figure 1 illustrates the electrophoretic mobilities of the albumin, alpha1, alpha2, beta and gamma protein bands on QuickGel Split Beta SPE Gel. The fastest moving band, and normally the most prominent, is the albumin band found closest to the anodic edge of the gel. The faint band next to this is alpha1, followed by alpha2, beta and gamma globulins.



**Figure 1:** A QuickGel Split Beta SPE Gel showing relative position of the bands.



**Figure 2:** A scan of a QuickGel Split Beta SPE pattern.

**Calculations of the Unknown**

The Helena QuickScan Touch Plus will automatically calculate and print the relative percent and the absolute value of each band when the total protein is entered. Refer to the Operator’s Manual provided with the instrument.

**INTERPRETATION OF RESULTS5**

Results on normal individuals will cover age and sex-related variations and day-to-day biologic variations. Disease states in which abnormal patterns are observed include inflammatory response, rheumatic disease, liver diseases, protein-loss disorders, plasma cell dyscrasias, pregnancy and genetic deficiencies.

**Further Testing Required**

The serum protein electropherogram or densitometric tracing should be evaluated for abnormalities. If abnormalities are observed, appropriate follow-up studies should be initiated. These may include immunoelectrophoresis, immunofixation, quantitation of immunoglobulins, bone marrow examination and other appropriate tests.

**LIMITATIONS**

Since all electrophoretic procedures are nonlinear, it is critical to fill the wells with the recommended volume of undiluted serum to obtain optimal resolution and reproducible results. Noncompliance with the recommended procedure may affect the results.

**SPECIFIC PERFORMANCE CHARACTERISTICS for spife  
PRECISION**

**Within Run:** A normal patient sample, a normal control and an abnormal control were run alternately on a single gel with the following results:

Normal Control N = 7

Protein Fraction Mean % SD CV\_\_\_

Albumin 57.1 1.1 2.0%

Alpha1 3.2 0.2 5.8%

Alpha2 10.1 0.4 3.9%

Beta 16.3 0.3 1.9%

Gamma 13.2 0.7 5.4%

Abnormal Control N = 7

Protein Fraction Mean % SD CV\_\_\_\_

Albumin 45.5 1.0 2.1%

Alpha1 3.1 0.2 7.9%

Alpha2 9.9 0.4 3.9%

Beta 12.7 0.3 2.1%

Gamma 28.7 0.3 1.1%

Normal Patient N = 6

Protein Fraction Mean % SD CV\_\_\_

Albumin 52.0 0.7 1.3%

Alpha1 3.3 0.1 3.6%

Alpha2 10.5 0.4 3.7%

Beta 15.8 0.2 1.2%

Gamma 18.4 0.4 2.0%

**Between-Run:** A normal patient sample, a normal control and an abnormal control were run in replicate on eight gels with the following results:

Normal Control N = 56

Protein Fraction Mean % SD CV\_\_\_\_

Albumin 56.4 0.9 1.5%

Alpha1 3.6 0.3 9.1%

Alpha2 10.3 0.4 3.4%

Beta 16.3 0.3 2.1%

Gamma 13.3 0.4 3.0%

Abnormal Control N = 56

Protein Fraction Mean % SD CV\_\_\_\_

Albumin 45.2 0.8 1.7%

Alpha1 3.5 0.3 8.8%

Alpha2 10.0 0.3 2.9%

Beta 12.7 0.3 2.1%

Gamma 28.7 0.4 1.4%

Normal Patient N = 48

Protein Fraction Mean % SD CV\_\_\_

Albumin 51.0 0.8 1.6%

Alpha1 3.7 0.2 6.7%

Alpha2 10.8 0.3 3.1%

Beta 16.0 0.3 2.1%

Gamma 18.4 0.4 1.9%

**Correlation**

Normal (N = 50) and abnormal (N = 50) serum samples were analyzed using the SPIFE Split Beta SPE system and the QuickGel Split Beta SPE system.

N = 100

Y = 0.962X + 0.619

R = 0.998

X = SPIFE Split Beta SPE

Y = QuickGel Split Beta SPE on SPIFE

**PERFORMANCE CHARACTERISTICS FOR QUICKGEL CHAMBER PRECISION**

**Within Run:** A normal patient sample, a normal control and an abnormal control were run alternately on a single gel with the following results:

Normal Control N = 7

Protein Fraction Mean % SD CV\_\_\_

Albumin 57.4 0.9 1.6%

Alpha1 3.4 0.2 5.5%

Alpha2 9.7 0.3 3.6%

Beta 18.2 0.5 2.6%

Gamma 11.3 0.5 2.6%

Abnormal Control N = 7

Protein Fraction Mean % SD CV\_\_\_\_

Albumin 53.9 0.8 1.4%

Alpha1 4.1 0.2 5.4%

Alpha2 11.3 0.3 2.3%

Beta 13.5 0.3 2.6%

Gamma 17.2 0.3 1.8%

Normal Patient N = 6

Protein Fraction Mean % SD CV\_\_\_

Albumin 57.9 1.0 1.7%

Alpha1 3.1 0.1 4.7%

Alpha2 9.4 0.3 3.3%

Beta 16.8 0.3 2.1%

Gamma 12.7 0.4 3.4%

**Between-Run:** A normal patient sample, a normal control and an abnormal control were run in replicate on four gels with the following results:

Normal Control N = 28

Protein Fraction Mean % SD CV\_\_\_\_

Albumin 58.2 1.3 2.3%

Alpha1 3.4 0.2 6.5%

Alpha2 9.4 0.5 4.8%

Beta 17.8 0.6 3.3%

Gamma 11.1 0.5 4.6%

Abnormal Control N = 28

Protein Fraction Mean % SD CV\_\_\_\_

Albumin 52.5 1.3 2.5%

Alpha1 4.3 0.4 5.5%

Alpha2 11.5 0.4 3.7%

Beta 13.8 0.5 3.7%

Gamma 17.9 0.5 3.3%

Normal Patient N = 24

Protein Fraction Mean % SD CV\_\_\_

Albumin 58.4 1.2 2.2%

Alpha1 3.1 0.2 5.8%

Alpha2 9.3 0.4 4.2%

Beta 9.3 0.4 4.2%

Gamma 12.4 0.7 5.3%

**Correlation**

Normal (N = 25) and abnormal (N = 25) serum samples were analyzed running the QuickGel Split Beta SPE Kit on the QuickGel Chamber and on the SPIFE 3000.

N = 50

Y = 0.957X + 0.853

R = 0.998

X = QuickGel Split Beta SPE on the SPIFE

Y = QuickGel Split Beta SPE on the QuickGel Chamber

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