Sperm Preparation for Intrauterine Insemination Using Density Gradient Separation

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1 Introduction

Sperm washing is performed to remove seminal plasma and increase sperm density prior to intrauterine insemination (IUI) [1]. Gradient sperm washing is highly effective at isolating motile sperm from the ejaculate of most men. IUI may be indicated in cases of poor postcoital testing due to hostile or absent cervical mucus, oligospermia, or the presence of sperm antibodies [2]. It may be used in conjunction with artificial insemination or IVF procedures.

2 Specimen Collection

The patient is instructed on how to collect the specimen (see “Semen Collection and Labeling Procedure”). The patient collects the specimen into a sterile container and brings it to the laboratory. The patient, or partner, is told to return in ~1 h to pick up the washed specimen.

3 Equipment and Materials

A. Components of the sperm separation kit—sterile colloidal suspension of silica particles stabilized with covalently bound hydrophilic silane supplied with HEPES-buffered human tubal fluid (HTF):
   1. Lower phase (80 %)
   2. Upper phase (40 %)

B. Sperm washing media (modified HTF with 5 % human serum albumin and gentamicin)
C. Disposable sterile 15 mL polystyrene conical centrifuge tube(s) with cap(s)
D. Centrifuge
E. 37 °C incubator
F. Sterile graduated serological pipettes (2 mL)
G. Computer-assisted semen analyzer (CASA)
H. Long stemmed (9”) pasteure pipette
I. Disposable 20 μm sperm counting chamber
J. Makler counting chamber
K. Dilution cups 2-mL
L. Brown paper bags
M. Disposable sterile transfer pipettes
N. Viscosity treatment system (when applicable)-5 mg chymotrypsin

4 Quality Control

A. Sperm recovery and percent motility are checked and recorded semiannually on all IUI samples.

   Criteria: None defined.

B. Daily Precision:

   A patient specimen should be selected at random and run through CASA. A manual count and motility reading should be performed simultaneously.

   Criteria: All manual results should be within 20 % of CASA value.

   Response: If results are not within the defined percentage difference, the sample must be repeated. If it is still out-of-range, inform supervisor or director.

C. Technologists review patient results to check for technical and clerical errors prior to release of specimen for insemination.
5  Procedure

5.1  Prepare Reagents

1. Bring the upper and lower phase, HTF, and semen sample to 37 °C by incubating for 20 min (Fig. 14.1).
2. Transfer 2 mL of the lower phase into a sterile disposable 15 mL conical centrifuge tube.
3. Layer 2 mL of the upper phase on top of the 2 mL lower phase using a sterile pipette. Slowly dispense the upper phase lifting the pipette up the side of the tube as the level of the upper phase rises.

Note: A distinct line separating the two layers should be observed. This two-layer gradient is stable for up to 2 h.

5.2  Prepare Paperwork/Accept Patient Specimen

1. Make sure the specimen container is labeled with two identifi cers. Acceptable identifi cers are either the patient name and date of birth or the patient name and medical record number (or SSN).

Note: If all of the patient information on the specimen cup is not present, the container should be labeled in front of the patient.
2. Have the “Sperm Wash Worksheet,” “Artifi cial Insemination by Husband,” “Specimen Drop-off/Pick-up Form,” and two IUI labels prefilled before the patient arrives with all pertinent information.
3. Label a 15 mL centrifuge tube with the patient’s and partner’s name, medical record numbers, and date. The tube should also be labeled with color-coded tape* as an extra identifi er.

5.3  Wash/Analyze Specimen

Note: Always use sterile technique during specimen processing.
1. Allow the semen specimen to liquefy completely for ~20 min in a 37 °C incubator before processing.

Note: Occasionally, samples do not liquefy properly and remain too viscous to pass through the gradient. Add 5 mg of chymotrypsin to the viscous sample ~5–10 min before layering to increase motile sperm yields.
2. Using a sterile transfer pipette, place a few drops of the patient’s semen sample into a conical cup. Place ~5 μL of the sample onto a sperm counting chamber and microscopically verify that there is motile sperm before proceeding with the wash (this sample can also be used for the prewash analysis).

*Note: Color-coded labeling tape should be used for all tubes, media, and paperwork and should be specific (of the same color) for each patient.
4. Have the person delivering the specimen sign the “Specimen Drop-off/Pick-up Form.” They must at this time present an acceptable ID card (e.g., driver’s license, employee badge). The technologist accepting the specimen must record on the sheet the type of identifi cation presented and the appropriate ID number.
5. Label a 2 mL conical cup for the post-wash analysis. Remove the warm tube of sperm wash media (HTF with 5 % HSA) from the 37 °C incubator and label it with the patient’s name and colored labeling tape* (same color as used above).

Fig. 14.1  Incubator set at 37 °C and depiction of sample undergoing liquefaction [Reprinted with permission, Cleveland Clinic Center for Medical Art & Photography © 2015. All Rights Reserved]
3. Measure the total semen volume using a sterile 2 mL serological pipette.

4. Gently layer no more than 3 mL* of liquefied semen onto the upper phase.
   *Note: If the sample volume is greater than 3 mL, it is necessary to split the specimen into two tubes for processing. Process the second tube using the same procedure that was used for the first tube. Combine the second tube with the first tube during step #11 after adding HTF media.

5. Centrifuge for 20 min at 1600 rpm (Fig. 14.1).

6. While the specimen is in the centrifuge, perform a pre-wash CASA analysis. Perform CASA analysis according to the “Routine Semen Analysis” protocol and record the results on the worksheet.
   Note: While examining the specimen, pay particular attention to any extraneous round cells, debris, and bacteria that are present. If round cells are greater than or equal to 1.00 M/mL, perform an Endtz test immediately. A positive Endtz test should be reported to the appropriate medical personnel as soon as possible. The Endtz result should also be written on the tube’s outer label at the end of the wash procedure before handing the specimen to the patient.

7. After the specimen has centrifuged, remove the supernatant without disturbing the pellet. Using a sterile long Pasteure pipette carefully remove the seminal plasma, upper interphase (raft “a”), upper (40%) colloid layer, and the lower interphase “raft b”). Leave the majority of the 80% colloid layer in place and discard the aspirated material.

8. Using another clean Pasteure pipette remove the soft pellet by direct aspiration (maximum 0.5 mL) from the bottom of the tube beneath the lower (80%) colloid layer into the soft. Avoid contaminating the pasteure tip aperture with the residual seminal plasma/raft material.

9. Transfer the pellet to a clean 15 mL conical tube.

10. Using a serological pipette, add 2 mL of HTF media and resuspend the pellet by pipetting the sample gently up and down until the sperm pellet is properly mixed into the sample.

11. Centrifuge the resuspended sample for 7 min at 1600 rpm (Fig. 14.2).

12. Repeat step 7 above (Fig. 14.3).

13. Resuspend the final pellet (Fig. 14.4) using 0.5 mL of HTF media with a 2 mL serological pipette. Note the volume of the sample. Leave a small droplet (~20 μL) in a labeled conical cup for the post-wash analysis. Subtract this volume from the volume noted above and record on the patient worksheet (Fig. 14.5).

14. Seal the tube using tamper-evident tape, label it with the premade label, and place it in the incubator until the patient arrives. Show the color-coded tube to the patient and have the patient verify the couple’s names, medical record numbers, and date. Place the sample in a brown paper bag and hand it to the patient. Be sure the patient signs the specimen pick-up form and presents appropriate identification. The patient should carry the specimen to gynecologist for the insemination. The gynecologist determines whether or not the sperm count and motility are sufficient for

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Fig. 14.2  Double-density gradient wash procedure; separation of seminal plasma, abnormal nonmotile sperm, and viable motile sperm [Reprinted with permission, Cleveland Clinic Center for Medical Art & Photography © 2015. All Rights Reserved]
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insemination. If no motile sperm are found on the post-wash specimen, page the gynecology nurse prior to releasing specimen.

**Note:** Always look at the post-wash specimen on a sperm counting chamber before releasing it to the patient. Notify the inseminator immediately if bacteria are present. If there is a positive Endtz test, report the result to the nurse as soon as possible and write it on the outside label of the tube before handing it to the patient. If the prewash Endtz is positive and the post-wash round cell count is less than 1 M/mL, note the round cell count on the outside label of the tube.

13. Perform a post-wash CASA analysis according to the “Routine Semen Analysis” protocol and record the results on the patient worksheet. Page the nurse with the patient name, clinic number, total motile sperm, Endtz value (if necessary), and any other pertinent information.

14. Record appropriate information (below) on the “Artificial Insemination by Husband” form:
   (a) MRN
   (b) Date of insemination
   (c) Total motile post-wash sperm
   (d) Wash type (e.g., gradient)
   (e) Insemination performed (Gyn nurse)
   (f) Comments (e.g., positive Endtz, agglutination, high viscosity)
   (g) Tech initials

15. After checking for technical and clerical errors, enter the results and print the final report. Check the final report to ensure the accuracy of the final results.

**Note:** Immediately correct any inaccurate results.

Fig. 14.3  HTF resuspended sample centrifuged to produce viable sperm pellet [Reprinted with permission, Cleveland Clinic Center for Medical Art & Photography © 2015. All Rights Reserved]
6 Procedural Notes

A. It is important to use sterile plasticware and glassware during the sperm wash procedure.
B. Each patient specimen should be kept in a separate specimen rack during the time of processing.
C. Processing of patient specimen(s) should be started and completed by a single technologist. In rare cases, when a second technologist is called to help during the processing, the patient’s paperwork should be reverified by the assisting technologist.

7 Tips on How to Maximize the Yield from Density Gradient

A. Make sure you have all components of the gradient brought to 37 °C temperature before use. Besides avoiding a “cold shock” to the spermatozoa, condensation on the media bottles disappears allowing for better inspection of the bottles for contamination. Any bottle whose contents appear in any way cloudy or hazy should not be used.
B. Do not use the same pipette in more than one bottle of media.
C. Density gradient components are air-buffered. Prolonged exposure to a 5 % CO$_2$ environment will alter the pH of these products, which may in turn affect their nature and performance.
D. For highly viscous semen add 5 mg of chymotrypsin to the ejaculate 10 min before it is placed on top of the gradient increases motile sperm yields without any measurable damage to the motile sperm.
E. Do not overload the gradient. The volume of semen in this case is less important than the absolute number of sperm placed on the gradient. Usually, a maximum number should be around 120 million cells. Overloading the gradient will result in a phenomenon called “rafting”—this aggregation of desirable, as well as undesirable, components of the semen will be
created and will be present in the post-centrifugation pellet.

F. Always use the gradient within 1 h after creating it—eventually the two phases over time blend into each other and a sharp interface will not exist.

G. To optimize the number of motile sperm, avoid contamination with the seminal plasma raft “a” or raft “b”, or the upper and lower layers.

H. Use a long stem preferably a 9” Pasteure pipette to carefully aspirate the soft pellet at the bottom of the tube beneath the 80% layer.

I. Transfer the final soft pellet into a fresh 15 mL conical tube for second step of centrifugation.

References


Sperm Preparation for Intrauterine Insemination

Procedure

Sperm preparation by density gradient method is highly effective at isolating mobile sperm for intrauterine insemination.

A. Preparation of Reagents

1. Bring the upper and lower phase gradients, HTF, and semen sample to 37°C by incubating for 20 minutes (Figure 1).

2. Transfer 2mL of the lower phase into a sterile disposable 15mL conical centrifuge tube.

3. Layer 2mL of upper phase on top of the 2mL lower phase using a sterile pipette. Slowly dispense the upper phase lifting the pipette to the side of the tube as the level of the upper phase rises.

Note: A distinct line separating the two layers should be observed. This two-layer gradient is stable for up to 2 hours.

Figure 1. Incubator set at 37°C and depiction of sample undergoing liquidation

B. Wash/Analyze Specimen

Note: Always use sterile technique during specimen processing.

1. Allow the semen specimen to liquify completely for ~20 minutes in a 37°C incubator before processing.

Note: Add 5 mg of chymotrypsin to the viscous sample ~5-10 minutes before layering to increase motile sperm yield.

2. Using a sterile transfer pipette, place a few drops of the patient's semen sample into a conical cup. Place ~5uL of the sample onto a counting chamber and microscopically verify the presence of mobile sperm before proceeding with the wash (this sample can also be used for the prewash analysis).

3. Measure the total semen volume using a sterile 2mL serological pipette.

4. Gently layer no more than 3mL of liquefied semen onto the upper phase (Figure 2).

Note: If the sample volume is greater than 3mL, it is necessary to split the specimen into two tubes for processing. Process the second tube using the same procedure that was used for the first tube. Combine the second tube with the first during step #12 after adding Sperm Wash Medium.

5. Centrifuge for 20 minutes at 1600 rpm (Figure 2 and 3).

Figure 2. Double-density gradient wash procedure; separation of seminal plasma, abnormal non-mobile sperm and viable mobile sperm.

6. While the specimen is in the centrifuge, conduct a pre-wash CASA analysis.

7. After the specimen has centrifuged, remove the clear seminal plasma, the upper (40%) interphase “raft”, upper (40%) colloid layer, and the lower interphase “raft”. Leave most of the 80% (colloidal) in place. Discard the aspirated material without disturbing the lower phase of the pellet. Removal of the supernatant should be done with a long stem (9") transfer or Pasteur pipette and the tip of the pipette should be kept just below the meniscus, away from the top of the pellet.

8. Use a sterile long Pasteur pipette to aspirate the soft pellet (Figure 3).

9. Transfer the soft pellet into a fresh 15-mL conical centrifuge tube.

10. Using a serological pipette, add 2mL of Sperm Wash Medium and resuspend the pellet by pipetting the sample gently up and down until the sperm pellet is properly mixed into the sample (Figure 4).

Figure 3. Sperm Wash Medium resuspended sample centrifuged to produce viable sperm pellet.

11. Repeat centrifugation at 1600 rpm for 7 minutes. Carefully remove the sperm wash medium (Figure 4). (Figure 3).

Figure 4. Removal of Sperm Wash Medium.

12. Resuspend the final pellet (Figure 5) using 0.5mL of Sperm Wash Medium with a 2mL serological pipette. Note the volume of the sample. Leave a small droplet (~20 µL) in a labeled conical cup for the post-wash analysis. Subtract this volume from the volume noted above and record on the patient worksheet.

Figure 5. Sperm pellet resuspended in Sperm Wash Medium.

13. Seal the tube using tamper-evident tape and label it with the pre-made label. Place it in the incubator until the patient arrives.

14. Perform post-wash CASA semen analysis according to the "Routine Semen Analysis" protocol.