RULES AND REGULATIONS FOR IN-VITRO FERTILIZATION
PURSUANT TO ACT 920 OF 1991

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AUTHORITY

The following rules and regulation’s for facilities for in-vitro fertilization in Arkansas are duly adopted and promulgated by the Arkansas State Board of Health pursuant to the authority expressly conferred by the laws of the State of Arkansas in Act 920 or 1991. They will further act to amend the Arkansas Code of 1987 and implement and augment Arkansas Insurance Commission Rules and Regulation pursuant to authority of Acts 779 and 268 of 1987.

PURPOSE

These rules and regulations have been prepared for the purpose of establishing criteria for minimum standards for the certification of facilities for in-vitro fertilization in Arkansas. By necessity, they are of regulatory nature, but are considered to be practical minimum design and operation standards for these facilities. These standards are not static and are subject to periodic revisions in the future as new knowledge and changes in procedure trends become apparent. However, it is expected that facilities will exceed these minimum requirements and that they will
not be dependent upon future revisions in these standards as a necessary prerequisite for improved services.

SECTION I. COMPLIANCE.

Any authorized representative of the Arkansas Department of Health shall have the right to enter upon or into the premises of any institution at any time in order to make whatever inspection is deemed necessary in accordance with the minimum standards and regulations prescribed herein.

SECTION II. RESPONSIBILITY.

All facilities for in-vitro fertilization shall have a organized Governing Body or, in its absence a person who shall be legally responsible for maintaining quality patient care, establishing policies for the facility, and assuring the conduct of the facility. The responsible party shall ensure that the facility:

A. Maintains proper standards of professional work in the facility;
B. Formulates policies and procedures pertaining to the operation of the facility;
C. Credentials and appoints members to the medical staff and other authorized staff;
D. Develops and maintains an ongoing, comprehensive quality assurance program which will monitor and resolve identified problems.

SECTION III. POLICY AND PROCEDURE MANUAL.

The Policy and Procedure Manual shall include at a minimum:

A. Detailed job descriptions and duties, by job title, of each employee;
B. Policies and procedures for the general administration of the facility and for each department, section, or service of the facility;
C. Standards of care;
D. Procedures provided by the facility with detailed information of each;
E. Orientation practices for new employees;
F. Infection control measures;
G. Cleaning, disinfecting, and sterilizing procedures;
H. Fire safety and other safety and disaster practices;
I. Proper routine methods of handling and storing any flammable or explosive agents;
J. Documentation of those personnel who may perform procedures and give patient instruction.

All polices and procedure shall be reviewed annually and updated if necessary. The first page of the manual shall have the current review date and signature of the person conducting the annual review.

SECTION IV. CREDENTIAL FILES.

An individual file shall be maintained for each physician/para-professional practicing in the facility and shall include at least the following:
A. Specific delineation of privileges, requested and granted;
B. A signed application with documentation of the applicant’s agreement to abide by the facility’s Bylaws, Rules and Regulations;
C. Verification of current Arkansas licensure;
D. Verification of at least three (3) references;
E. Documentation of all actions taken concerning the applicant and indicating the types of the privileges granted and other applicable data.

SECTION V. QUALITY ASSURANCE.

There shall be an ongoing, comprehensive Quality Assurance program established for each facility which will monitor and resolve any identified problems. The program will include a written plan which shall include:

A. Methods of evaluating all services within the facility;
B. The evaluation of applicable procedures;
C. The evaluation of infection control for the facility.

There must be documented evidence to indicate that the facility has taken appropriate remedial action to address deficiencies identified through the quality assurance program. There shall also be documented evidence of appropriate action taken regarding each of the problems identified through the quality assurance program. Monitoring of the problem should continue until the problem is completely resolved with documentation of significant findings, conclusions, recommended actions, and results of actions.

SECTION VI. ADMINISTRATION.

A. All medical, surgical laboratory, and procedural care rendered in a facility for in-vitro fertilization certified by the Arkansas Department of Health shall conform to the current standards acceptable to the American Medical Association, the American College of Obstetricians and Gynecologist, the American Fertility Society, and the Clinical Laboratory Improvement Amendments of 1988 (CLIA ’88).

B. All containers used in the facility shall be legible and accurately labeled as to content.

C. There shall be a written disaster plan which shall include provisions for complete evacuation of the facility. The disaster plan should be rehearsed annually with written reports and evaluations of all drills maintained.

D. Fire extinguishers of the correct type shall be provided in adequate numbers, and shall be properly located and installed. Personnel shall be trained in the proper use of the equipment and in the procedures for proper fire containment and evacuation of the facility.

E. All refrigerated areas, including freezers, shall be provided with reliable thermometers, and records shall be maintained to document the temperatures on a daily basis or at least on each day of operation of the facility.

F. The facility must have a written plan covering collection, handling, and disposal of all types of waste including toxic, infectious, and non-infectious.
G. Ultrasonography services must be available to the facility seven (7) days per week.

SECTION VII. PERSONNEL.

A. The facility shall maintain a sufficient number of personnel to provide effective patient care and all other related services to an in-vitro fertilization program. There shall be written personnel policies and procedures which shall be made available to personnel. Provisions shall be made for orientation and continuing education programs for personnel.

B. Each employee shall have a health file. The file shall document at least:
   1. Annual testing for tuberculosis;
   2. Documentation of other tests required by the facility;
   3. Certification of current training in CPR techniques for all personnel involved in surgical/invasive procedures.

C. Written job descriptions shall be developed for each employee classification and shall include as a minimum the responsibilities or actual work to be performed in each classification. In addition, the job descriptions shall include the physical, educational, and licensing or certification requirements for each job classification. These job descriptions shall be included in the Policy and Procedure Manual for the facility.

D. Personnel records shall be maintained for each employee and shall contain no less than:
   1. Current and background information covering qualifications for employment;
   2. Records of all required health examinations;
   3. Evidence of current registration, verification, or licensure of personnel subject to statutory regulation.

E. An in-vitro fertilization program must include, as a minimum, personnel with the following expertise. A single individual may fulfill the requirement for expertise in one (1) or more areas.
   1. An individual with training and experience in reproductive endocrinology, particularly in the use of ovulation-inducing agents and the hormonal control of the menstrual cycle. The Arkansas Department of Health would accept an individual who has completed a Board-approved Fellowship in Reproductive Endocrinology for fulfillment of this requirement.
   2. An individual with expertise in pelvic reparative (infertility) surgery, as well as experience in laparoscopic and ultrasound-guided oocyte retrieval techniques. This combined expertise is necessary not only for the performance of the oocyte retrieval procedures, but also to ensure that the infertile couple is offered the most appropriate treatment modality.
   3. A director of the embryology laboratory. (See also Embryology Laboratory Personnel section.) This laboratory director must have personal experience in the organization and maintenance of a basic or clinical embryology laboratory, as well as in tissue culture techniques.
   4. An ultrasonographer (or obstetrician-gynecologist with specialized training and
experience in gynecologic sonography) who provides the monitoring of follicular
development and supervises ultrasound-directed oocyte retrieval.

5. Each program must have a designated overall program director. If the overall
program director is not a licensed physician, there must be designated a Medical
Director who is responsible for the clinical aspects of the treatment program.

6. An individual experienced in male reproduction (andrology) with special
competence in semenology. If this individual is not a urologist, then there should be
available consultant urologist services with expertise in reproductive surgery. This
individual should also be involved, when indicated, in recommending the most
appropriate treatment options for the couple.

7. In addition to the laboratory director, there should be a minimum of one (1) other
embryology technologist. Every embryologist should have a working knowledge of
sterile technique and cell tissue culture.

8. If gamete and/or embryo cryopreservation if offered, there should be an
individual with specialized training in cryobiology and experience in gamete and
cryopreservation techniques.

9. If oocyte micromanipulation is offered, there should be an individual with
specialized training in gamete biology and experience in micromanipulation techniques.

SECTION VIII. MEDICAL RECORDS.

A. A medical record shall be maintained for each patient of the in-vitro fertilization facility.
All these records must be kept separate from the other regular records of the facility. The
original or a copy of the original (when the original is not available) of all reports shall be filed
in the medical record. The record shall be permanent and shall be either typewritten or legible
written in ink. All dictated reports shall include the date of dictation and the date of
transcription. Only standard abbreviations, approved by the staff of the facility, shall be used.
This list of abbreviations shall be reviewed annually and revised if necessary.

B. Medical orders. All medical orders (medication, treatments, tests, and procedures) shall
be in writing and shall be signed by the physician.

C. Confidentiality. Medical records shall be considered confidential. Only authorized
personnel shall have access to the medical records. All medical records shall be secured at all
times; if authorized personnel are not present, the records shall be locked. Records shall be
available to authorized personnel from the Arkansas Department of Health.

D. Consent for Procedures. A specific consent for procedures shall be documented prior to
the procedure to be performed and shall include the date, time, and signatures of the patient, the
physician, and a witness.

E. A History and physical examination shall be documented prior to the procedure.

F. Anesthesia. When anesthesia is utilized, with or without loss of consciousness, a
complete anesthesia report, including pre-evaluation and post-follow-up shall be documented by
the Anesthesiologist and/or the Certified Registered Nurse Anesthesiologist (CRNA) or by the
Physician who administered the anesthesia.
G. Procedure Report. An individualized Procedure Report shall be written or dictated by the physician immediately following the procedure and shall be signed within seventy-two (72) hours. The report shall describe (in detail) techniques, findings, pre- and post-procedural diagnosis, and other pertinent information.

H. The record of the patient shall also include:
   1. Orders and reports of diagnostic services;
   2. Documentation of any medication administered;
   3. Progress notes for subsequent clinic visits recorded by applicable disciplines.

SECTION IX. UNITED STATE IN-VITRO FERTILIZATION REGISTRY (U.S. IVF REGISTRY) (S.A.R.T.).

A. In view of the continuing controversy about the success rate of in-vitro fertilization, all facilities performing this procedure should participate in the United State In-Vitro Fertilization Registry (U.S. IVF Registry) (S.A.R.T.). Furthermore, it is recommended that each program release (or permit the release from the Registry) identifiable clinic-specific success rates in order that patients and physicians may make appropriate choices among programs. As a minimum, the release data shall include the number of stimulation cycles begun, the number of oocyte retrieval procedures attempted, the number of women who became pregnant, and most importantly, the number of women who deliver live babies. Annualized statistics should be available within twelve (12) months of the completion of the year’s treatment cycles in order that the outcome of all the established pregnancies are available for inclusion.

B. In-vitro fertilization programs that choose not to participate in the U.S. IVF Registry shall prepare similar summaries of their results.

C. Records shall be kept of all attempts, with both successes and failures documented. Summaries of all records shall be available of correlation. The records shall include:
   1. Oocyte recovery;
   2. Fertilization;
   3. Cleavage;
   4. Conceptus transfer;
   5. Biophysical monitoring and fetal growth;
   6. Pregnancy outcome;
   7. All complications;

SECTION X. REQUIRED REASONABLE SUCCESS RATE.

There shall be a required reasonable success rate as defined by current American Fertility Society standards.

SECTION XI. INFORMED CONSENT.

A. As with all medical procedures and treatments, the patients must make the final decision
on what is appropriate and acceptable treatment in their particular situation. To comply with this requirement, it is necessary that each prospective patient couple be provided with full disclosure, including as a minimum, the program’s own experience with the specific procedure in question, including:

1. How long the facility has been performing it;
2. How many times the facility has performed it;
3. What the facility’s past and current success rates are.

B. This informed consent requirement shall be met with either the facility’s summaries or the information for the U.S. IVF Registry.

C. In addition to the provided information, in-vitro fertilization programs shall provide to couples full information concerning alternative procedures available to circumvent their specific infertility problem, including procedures that are not performed by the treating facility. Attention should also be given to the emotional needs and to the support of these patient couples.

SECTION XII. MEDICATIONS.

A. Security. All medications shall be locked when authorized personnel are not present.

B. Crash carts or emergency medication kits shall be provided. These carts/kits shall be secured with a breakaway seal and stored in an area observable by licensed personnel. The cart/kit shall have affixed a listing of the contents.

C. Refrigeration. Refrigeration shall be provided for the proper storage of biological and other drugs. An internal thermometer shall be provided and routinely checked to assure temperatures between 36 and 46 degrees Fahrenheit.

D. Controlled Substances.

1. Records. All controlled substances shall be stored to comply with State and Federal regulations. Records of disposition of controlled substances shall be maintained. The records shall include:
   a. Actual dosage administered to the patient;
   b. The patient’s name;
   c. Name of the physician;
   d. The date, time, and signature with title of the person administering the medication.

2. Breakage or Wastage. When breakage or wastage of a controlled substance occurs, the amount given and the amount wasted must be recorded by the licensed person who wasted the drug and verified by the signature of a licensed person who witnessed the wastage, and how it was wasted.

3. Inventory. There shall be a count at the opening and closing of the facility of all controlled substances stocked in the facility. The counts shall be made by two (2) licensed personnel (one (1) to count and one (1) to witness), both of whom shall sign the record with a notation made as to date and time of such count. Discrepancies in the count
are to be noted and reported to appropriate personnel.

E. **External Use Only.** Solutions and medications for external use only shall be kept separate from other medication.

F. **Errors and Adverse Drug Reactions.** Medication errors and adverse drug reactions should be documented and reported to the attending physician.

G. **Proper Maintenance ALL Drugs.** Drug storage areas should be routinely checked to ensure proper stock levels, drugs are in date, and containers are properly labeled, stored, and intact.

**SECTION XIII. SURGICAL SERVICES.**

A. Operating room facilities shall be provided for laparoscopic retrieval and ultrasound-guided retrieval.

B. The operating room availability must be rigidly controlled so patients can be moved in and out seven (7) days a week with a maximum of twenty-four (24) hours notice.

C. Anesthesia must be available seven (7) days a week.

D. A facility of oocyte and sperm culturing shall be close to the operating room with two (2)-way communication.

E. Any in-vitro fertilization facility offering services on an outpatient basis that require the use of general or intravenous anesthetics that render the patient unable to be responsible for his/her actions, and where, in the opinion of the attending physician, hospitalization is not necessary, shall comply with all applicable Sections 0100 through 1800 (excluding 0900 and 1200 through 1600) of Rules and Regulations for Hospitals and Related Institutions for Hospitals and Related Institutions in Arkansas of the Arkansas Department of Health. The most current rules and regulations shall apply.

F. An in-vitro fertilization procedure is so complex and highly technical that a large back-up system of laboratory support and specialized personnel should be available seven (7) days a week.

G. Other assisted reproductive technology (ART) that can include, but not be limited to, in-vitro fertilization (IVF), gamete intra-fallopian transfer (GIFT), tubal embryo transfer (TET), zygote intra-fallopian transfer (ZIFT), and others should be considered to be governed by these rules and regulations. All facilities licensed or certified by the Arkansas Department of Health offering these services shall conform to the current guidelines of the American College of Obstetricians and Gynecologists and also meet the minimal standards of the American Fertility Society.

**SECTION XIV. EMBRYOLOGY LABORATORY PERSONNEL.**

A. **Embryology Laboratory Director.** The embryology laboratory director shall be an individual with demonstrated knowledge of all laboratory aspects of Assisted Reproductive Technology (ART). To be acceptable as an embryology laboratory director, an applicant must fulfill both the following requirements:

   1. Hold an earned doctorate degree (Ph.D.) from an accredited institution in a chemical, physical, or biological science as the major subject or an M.D. degree from an
accredited institution. The laboratory director should have expertise and/or special training in biochemistry, cell biology, and physiology of reproduction with experience in experimental design, statistics, and problem solving/trouble shooting. The laboratory director should be responsible for formulating laboratory policies and protocols and should communicate regularly with the medical director regarding patient progress and protocols as they affect the laboratory aspects of treatment.

2. Have two (2) years documented pertinent experience in a program performing IVF- or ART-related procedures. This experience should include:
   a. Familiarity with laboratory quality control, inspection, and accreditation procedures;
   b. Detailed knowledge of tissue culture, ART, and andrology procedures performed in mammalian systems.

B. Laboratory Supervisor.

1. Education. If the medical director is also the laboratory director, there should be a qualified, designated laboratory supervisor. The embryology laboratory supervisor should have a Bachelor’s or Master’s degree in a chemical, physical, or biological science.

2. Experience. The embryology laboratory director or supervisor should have had a period of training of at least six (6) months and at least sixty (60) completed ART procedures in a program that performs at least one-hundred (100) IVF procedures per year with a minimum annual ten percent (10%) IVF live birth rate per retrieval cycle. A procedure is defined as a combination of the examination of follicular aspirates, insemination, documentation of fertilization, and preparation for embryo transfer. Satisfactory completion of this period of training should be documented by a signed letter from the laboratory director of the training program. In lieu of formalized training, a similar experience with the director’s own program is sufficient, provided the program has performed at least one-hundred (100) total retrievals and has had at least an annual ten percent (10%) live birth rate per retrieval cycle.

C. Laboratory Technologists.

1. Education. The embryology laboratory technologist should have an earned Bachelor’s degree from an accredited institution with a physical, chemical, or biological science as the major subject. Individuals without this educational requirement are acceptable provided they meet all the requirements in the following paragraph by January 1, 1992.

2. Experience. The embryology laboratory technologists should have documented pertinent experience in tissue culture and sterile technique with evidence of completion of thirty (30) complete IVF procedures under continuous supervision of the laboratory director or supervisor. Experience and documented training in tissue culture, sperm-egg interaction, or related areas of animal reproduction are desirable. The embryology laboratory technologist works under the supervision of a laboratory director or supervisor. Programs for the appropriate training of embryology laboratory technologists should be in place for each program, with documentation of completion for each
D. **Laboratory Staff.**

1. **Quantity of Procedures.** Each staff embryologist (including the embryology laboratory director or supervisor) should perform at least twenty (20) complete ART procedures a year.

2. **Expertise.** Among the embryology laboratory staff there should be one (1) or more persons with knowledge and experience in the following fields:
   - a. Preimplantation embryology;
   - b. Andrology;
   - c. Pre- and post-fertilization events.

3. **Knowledge of freeze-thaw.** Before attempting human embryo freezing, at least one (1) member of the embryology laboratory staff should have frozen in excess of two-hundred-fifty (250) animal embryos and should have demonstrated the ability to freeze-thaw embryos with a survival/development rate of over fifty percent (50%).

E. **Consultant Visits.** If the laboratory director is not physically present on a routine basis, consultant visits will be allowed. These visits are to ensure compliance with all applicable laws and regulations as well as current standards of practice. The visits should also identify current problems, note trends, and make recommendations for correction and improvement. The visits should be bi-monthly at the least, while monthly visits are strongly recommended. Each visit should be documented with a consultant’s report. In addition, relevant inservice educational programs should be provided during each visit.

**SECTION XV. LABORATORY SERVICES.**

A. **CLIA.** The laboratory must meet the conditions set forth under the Clinical Laboratory Improvement Amendments of 1988 (CLIA ’88) in order to be certified to perform testing on human specimens. All laboratory services, including andrology and embryology, are subject to CLIA ’88. A referenced laboratory may be used.

B. Rapid hormonal assays of estradiol and luteinizing hormones must be available seven (7) days per week.

**SECTION XVI. PHYSICAL ENVIRONMENT OF OCYTE COLLECTION/PROCEDURE ROOM; EMBRYOLOGY LABORATORY; ANDROLOGY LABORATORY.**

A. **Size** of the oocyte collection/procedure room must meet all of the requirements of a general operating room, with a minimum clear area of two-hundred-fifty (250) square feet exclusive of fixed and movable cabinets and shelves.

B. **Air.** The air filtration system for the procedure room and for the embryology laboratory must provide twenty (20) air changes per hour. If the rooms are adjacent, there must be positive pressure from the embryology laboratory room to the procedure room.

C. **Temperature.** The temperature in the procedure room and in the embryology laboratory must be maintained between sixty-eight (68) and seventy-four (74) degrees Fahrenheit.
D. **Humidity.** The humidity in the procedure room and in the embryology laboratory must be maintained between thirty-five (35) and fifty (50) percent.

E. **Walls and Floors.** Walls and floors must be composed of materials which are easily washed and disinfected and that are not physically affected by germicidal and cleaning solutions. Wall bases must be made integral and covered with the floor, tightly sealed within the wall, and constructed without voids that can harbor insects.

F. **Ceilings.** Ceilings must have a smooth finish which is washable and waterproof.

G. **Adequate space and mutual accessibility.** The laboratory must be constructed and arranged to ensure adequate space for the performance and reporting of test and for equipment. The embryology laboratory must be located within one-hundred (100) feet of the oocyte collection/procedure room. The presence of a general access hallway between the embryology laboratory and the procedure room is not recommended. As an alternative, oocyte identification and isolation can be performed in the procedure room, using a self-contained, temperature- and environmental- controlled, microscope/incubator unit. Oocytes can then be transported within this unit (or in a portable incubator) to the embryology laboratory.

H. **Safety.**
   1. Aerosols and toxic pest control substances must not be used.
   2. Non-toxic (non-powdered) gloves must be worn while handling gametes and embryos.
   3. Toxic fumes which could be harmful to gametes and embryos must be eliminated.
   4. All compressed gas cylinders must be secured to prevent falling.

**SECTION XVII. INFECTION CONTROL.**

Universal precautions must be observed.

A. **Medical Waste.** The facility must comply with the Rules and Regulations Pertaining to the Management of Regulated Medical Waste from Health Care Related Facilities promulgated under the authority of Act 96 of 1913 as amended, and Act 41 of 1992.

B. **Testing for Communicable Disease.**
   1. Both partners must be tested for hepatitis B and for the human immunodeficiency virus (HIV) in the three (3) months prior to initiation of the ART treatment cycle.
   2. A full bacteriologic examination must be conducted if the initial semen analysis reveals any sign of infection.

C. Goggles or glasses and face masks as appropriate must be worn while handling gametes and embryos.

**SECTION XVIII. EQUIPMENT MAINTENANCE**

A. **Maintenance Records.**
   1. There must be a documented preventive maintenance program for each piece of equipment, based upon the manufacturer’s recommendations.
   2. Maintenance records must be maintained for the life of the instrument.
3. There must be documentation of remedial action taken when there is equipment failure.

B. **Individual usage records.** There must be documentation of the following on each day that the facility is open if the particular piece of equipment is in use:
   1. Temperature of each refrigerator, freezer, and incubator;
   2. Adequate humidity of each incubator;
   3. External verification of the gaseous environment in each incubator;
   4. Level of liquid nitrogen in any embryo or gamete storage container;
   5. Status of gas cylinders and/or liquid nitrogen reservoir supplies.

C. **Alarm Systems.** Incubators and frozen embryo storage facilities must be equipped with alarms which would sound at a location which is staffed twenty-four (24) hours per day. Alarms must be tested at least monthly for proper operation.

D. **Emergency Generator.** There must be an automatic emergency generator in the event of power failure. The system must be exercised weekly and tested under load monthly.

**SECTION XIX. QUALITY CONTROL.**

There must be a written quality control program, which includes control bioassays, designed to check cell culture media and contact materials for toxins, inappropriate ionic concentration, microbial contamination, or other potential hazards to human gametes or embryos.

A. **Documentation.**
   1. **Quality Control Procedures.** There must be documentation of all quality control procedures performed.
   2. **Remedial Action.** There must be documentation of remedial actions when control procedures do not meet the laboratory’s criteria for acceptability.
   3. **Culture Medium.** There must be documentation of the following on each batch of culture medium:
      a. Date prepared;
      b. Name of person who prepared the medium;
      c. Osmolarity;
      d. pH;
      e. Method of Sterilizations;
      f. Expiration;
      g. Results of control procedures.

B. All quality control records must be retained for a period of two (2) years.

**SECTION XX. EMBRYO CRYOPRESERVATION.**

A. **Records.** Duplicate records on frozen embryos must be maintained in the laboratory and
with the program director. Records must contain the following information:

1. Developmental stage at which frozen;
2. Freezing protocol used;
3. Recommended thawing procedures;
4. Physical location of each embryo within the storage container.

B. Embryo containers must be labeled with the following:

1. Patient’s name;
2. Patient identification number;
3. Embryo identification number;
4. Date of cryopreservation.

C. There must be written procedures for the verification of patient identity and the identification of gamete and embryo samples prior to embryo thawing.

SECTION XXI. REQUIRED EQUIPMENT.

A. In-vitro fertilization. The following equipment is essential to any in-vitro fertilization program:

1. Incubator;
2. Centrifuge;
3. Microscope;
4. Warming block or Isolette;
5. Laminar flow hood.

B. Cryopreservation. The following equipment is essential to any in-vitro fertilization program which perform cryopreservation:

1. Controlled rate freezer;
2. Liquid nitrogen storage container.

SECTION XXII. SEVERABILITY.

If any provision of these Rules and Regulations, or the application thereof to any person or circumstances is held invalid, such invalidity shall not affect other provisions or applications of these Rules and Regulations which can give effect without the invalid provisions or applications, and to this end the provisions hereto are declared to be severable.

SECTION XXIII. REPEAL.

All Regulations and parts of Regulations in conflict herewith are hereby repealed.

CERTIFICATION
This will certify that the foregoing Rules and Regulations Pertaining To In-Vitro Fertilization were adopted by the State Board of Health of Arkansas at a regular session of said Board held in Little Rock, Arkansas, on the 22nd day of July, 1993.

Tom S. Butler  
Secretary of Arkansas State Board of Health  
Acting Director, Arkansas Department of Health

Dated at Little Rock, Arkansas, this 24th day of September, 1993.

The foregoing Rules and Regulations, copy having been filed in my office, are hereby approved on this 30th day of September, 1993.

Jim Guy Tucker  
Governor