Simplified Embryo Vitrification Protocol for Cryolock
2PN to Blastocyst

ALL PROCEDURES MUST BE PERFORMED AT ROOM TEMPERATURE (22–27°C)

Prior to vitrification of blastocysts, refer to Collapsing Protocol, Human Blastocysts, FISI P/N 002107.

Have all necessary materials, tools, and equipment ready and easily accessible before starting procedure.

1. Aseptically dispense one (1) 50 µL drop of ES.
2. Transfer embryo(s) (2 maximum) to the ES drop and expose undisturbed for 6–10 minutes.
   - The specimen(s) will shrink and then gradually return to original size, indicating that equilibration is complete.
3. During above equilibration in ES, aseptically dispense one (1) 50 µL drop of VS 2 minutes prior to complete equilibration.
4. Transfer embryo(s) with minimal volume of medium from ES drop to the VS drop for 30 seconds before loading.
5. Gently but thoroughly pipette embryo(s) within VS drop to ensure complete rinse with VS.
   - To minimize floating, after 10 seconds pipette the specimen(s) to the bottom center of the VS drop.
6. Load, seal, and plunge Cryolock within 80 seconds, not to exceed 110 seconds after initial exposure to VS.
   - Load after specimen is completely dehydrated and stable at the bottom of the VS drop.
7. Refer to Cryolock loading protocol and product insert for detailed loading instructions and warnings.
   - See reverse side for tips.

KEY

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Simplified Embryo Vitrification Protocol for Cryolock
Tips

- All procedures are to be done at ROOM TEMPERATURE (22–27°C).
  Do not use heated stage.
- Have all necessary materials, tools, and equipment ready and easily accessible before starting procedure.
- Cryolock tip should be checked and Cryolock should be pre-labeled with patient information before starting.
- Where possible, select only the best quality embryos (2PN to Blastocyst) for vitrification.
- The recommended Cryolock capacity is a MAXIMUM of 2 specimens.
- Process only as many specimen(s) as will be loaded per Cryolock at one time.

- Minimize exposure of specimens to light during equilibration in ES and VS solutions.
- Transfer specimens between drops using a minimal volume of medium.
- The timing for exposure to VS is CRITICAL:
  - Maintain microscopic visualization of specimen(s) by adjusting focus as needed during rapid exposure to VS (specimens will float in the drop).
  - Keep transfer pipette tip close to drop for quick manipulations.
  - Load, seal, and plunge the Cryolock within 80 seconds, not to exceed 110 seconds after initial exposure to VS.

*Use of Cryolock with blastocysts is not cleared in the U.S.
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