Standard Operating Procedure for Preparation of *Aspergillus fumigatus* Test Strains for Inhalational Pulmonary Aspergillosis Animal Studies

1. Purpose
   This Standard Operating Procedure (SOP) will provide information necessary for the uniform completion of *Aspergillus fumigatus* inoculum preparations for use in infecting laboratory animals with experimental pulmonary aspergillosis.

2. Scope
   This SOP will provide sufficient information to allow preparation of a sufficient volume of concentrated *Aspergillus fumigatus* conidia for use in either the Madison or Acrylic inhalation chambers.

3. Definitions.
   “Inoculum preparation” will mean to create a precise, quantified concentration of viable *Aspergillus fumigatus* conidia in a diluent suitable for suspending and stabilizing the same.

4. Responsibilities
   This SOP shall be utilized by employees of Research assistant status or higher without additional training. Research technicians may perform this work upon receipt of training.

5. Equipment and Materials
   - **Method A. (preferred)**
     - Sterile phosphate buffered saline (PBS) supplemented with 0.1% v/v Tween 80
     - RC5B High Speed Centrifuge (Sorvall)
     - Sorvall SS-34 rotor
     - Sterile disposable plastic inoculation loops (Fisher)
     - Sterile Oakridge tubes (30 ml)
     - 50 ml sterile polypropylene centrifuge tubes with flat top rim seal caps
     - Parafilm
     - Potato dextrose agar plates (PDA)
     - Hemacytometer
     - Amphyl (Revco) [Acceptable equivalent: Vespheune (Steris) or Decon (Decon Labs)]
   - **Method B. (alternative)**
     - Sterile Phosphate buffered saline (PBS) with 0.1% v/v Tween 80
     - Beckman Model TJ-6 Table-top Centrifuge or equivalent
     - Sterile disposable plastic inoculation loops (Fisher)
     - Potato dextrose agar plates (PDA)
     - Parafilm
     - 50 ml sterile polypropylene centrifuge tubes with flat top rim seal caps
     - 5 ml sterile snap top polycarbonate tubes
Glass wool (2 x 2 piece) inserted into a 30 ml sterile glass syringe and sterilized as a unit (2)

Hemacytometer

Amphyl (Revco) [Acceptable equivalent: Vesphene (Steris) or Decon (Decon Labs)]

6. Procedure

**Preparation of inoculum for 1 chamber run**

- **Method A. (preferred)**
  - One chamber run will inoculate 70 mice using 10⁹ conidia/ml of *Aspergillus fumigatus* suspension or 9 guinea pigs using 7x10⁷ conidia/ml of *Aspergillus fumigatus* suspension.
  - All manipulations should be performed at room temperature in a laminar flow hood or biological safety cabinet, with disposable hospital pads lining the work area unless otherwise indicated.
  - Ten days before the planned day of infection inoculate PDA plates (10 plates will be needed per chamber run for mice. 6 plates will be needed for each chamber run for guinea pigs) for each planned run with 10 µl of test strain (i.e. *Aspergillus fumigatus* 293) conidial stock suspension pipetted to the center of the plate. Spread the inoculum all over the surface of the plate with a sterile loop. Seal inoculated PDA plates with parafilm, invert and grow at 37°C until the day of infection.
  - Note: Seven days prior to the planned day of infection, inoculate an additional 5 PDA plates with the test strain to ensure contaminant-free plates of the test strain are available for harvesting conidia in the event the first 10 are contaminated or there are problems with the growth of the strain.
  - On the day of infection harvest conidia with sterile PBS + 0.1% v/v Tween 80 (PBST). Scrape the entire plate with a sterile disposable plastic loop ensuring that you only scrape the surface and not dig into the agar or scrape too vigorously as to spill buffer outside of the plate and reduce hyphal contamination. Use 10 ml PBST per plate. Repeat a second time by pipetting and aspirating with 5 ml of PBST until conidia become suspended and the liquid is dark green.
  - Collect the PBST plus conidia in high speed Oakridge centrifuge tubes, and concentrate by high speed centrifugation (we use 15,000 rpm ['avg 17,000 X g] for 15 minutes, at 4°C no brake in a Sorvall SS-34 rotor).
  - After the spin, decant the supernatant from all tubes into a sterile 150 ml flask, leaving approximately 5 ml volume in one tube. Reserve the supernatant for re-spinning in case insufficient numbers of conidia were isolated. Resuspend the pellet in the 5 ml
volume by vortexing. If the supernatant is not needed, discard into a large beaker containing Amphil disinfectant.

- Decant the suspended conidia and add it to the second centrifuge tube containing only the conidial pellet and no supernatant. Rinse the first tube with 5 ml of the reserved supernatant, add to the second tube (total volume 10 ml) and resuspend the second pellet. Repeat as necessary until all conidia are in one tube or flask, which will vary based on the number of animals to inoculate, and therefore, the number of runs. Vortex vigorously but do not sonicate. Count a 1:1000 or 1:10,000 dilution in PBST with a hemacytometer for a target inoculum of $1 \times 10^9$ conidia/ml for mice or $7 \times 10^7$ conidia/ml for guinea pigs.

- Confirm inoculum viability by serially diluting an aliquot of the inoculum in PBST. Prepare serial dilutions of the stock, and plate 100 µl of $1 \times 10^{-5}$, $10^{-6}$ and $10^{-7}$ dilutions of conidia suspensions for mouse inoculation – or – plate 100 µl of $1 \times 10^{-4}$, $10^{-5}$ and $10^{-6}$ dilutions of conidia suspensions for guinea pig inoculation as appropriate. Incubate plates sealed with parafilm at 37°C overnight and count the colonies the next day.

- Conidial stock should be stored at 4°C until use.

**Method B. (alternative):**

- All manipulations should be performed at room temperature in a laminar flow hood or biological safety cabinet, with disposable hospital pads lining the work area unless otherwise indicated.

- Ten days before the planned day of infection inoculate PDA plates (40 plates will be needed per chamber run for mice and 20 plates will be needed for each chamber run for guinea pigs) for each planned run with 10 µl of test strain (i.e. *Aspergillus fumigatus* 293) conidial stock suspension pipetted to the center of the plate. Seal inoculated PDA plates with parafilm, invert, and grow at 37°C until the day of infection.

- Note: Seven days prior to the planned day of infection, inoculate an additional 10 PDA plates with the test strain to ensure contaminant-free plates of the test strain are available for harvesting conidia in the event the first 40 are contaminated or there are problems with the growth of the strain.

- On the day of infection harvest conidia with sterile PBS + 0.1% v/v Tween 80 (PBST). Scrape the entire plate with a sterile disposable plastic loop ensuring that you only scrape the surface and not dig into the agar or scrape too vigorously as to spill buffer outside of the plate and reduce hyphal contamination.

- Aspirate the 10 ml of PBST + conidia from the plate and decant into a 50 ml conical tube. Add an additional 5 ml of sterile PBST
to wash the surface of the plate and combine the washes in the same conical tube.

- Repeat as necessary for the number of plates at hand. A total of 30 plates will normally be needed to yield $10^9$ CFU/ml per one run of either the Madison or the acrylic chamber for mice. A total of 15 plates will normally be needed to yield $\sim 10^8$ CFU/ml per one run of either the Madison or the acrylic chamber for guinea pigs.

- Collect and concentrate conidia in 50 ml conical tubes by centrifugation (in Beckman Model TJ-6 Table-top Centrifuge with a Beckman TH-4 swing bucket rotor) at 2500 rpm \( [\text{avg } 1092 \times g] \) for 15 minutes at room temperature.

- Discard supernatant into a large beaker (containing amphi disinfectant) taking care not to disrupt pellet (the pellet will not be very compact). Leave about 5 ml of supernatant in one tube and vortex to re-suspend pellet. Repeat with the second tube and combine the re-suspended conidia with the contents of the initial tube. Wash tube 2 with 5 ml of sterile PBST and add this to tube 1 and discard tube 2. Repeat with subsequent tubes until tube 1 reaches a volume of 40 ml and start with a new tube. Repeat until the number of tubes are less than $\frac{1}{2}$ the number of what you started. (You may also use a large sterile flask to collect all the conidia and washes in one container at this time only).

- Place the autoclaved glass wool/syringe unit in a 50 ml conical tube. Pour the entire conidial solution through until the tube is filled (approximately 40 ml). Replace tube with a new one and repeat. (This is done to filter out any hyphal fragments that may be present). If necessary, replace the glass wool/glass syringe unit with another if the flow through begins to slow or is blocked.

- Concentrate conidia by centrifugation at conditions previously described and combine all pellets into one tube for one last centrifugation.

- Re-suspend the pellet in 30 ml of sterile PBST. Vortex vigorously.

- Make 1:10,000 and 1:100,000 dilutions of the conidial stock in sterile PBST.

- Count the dilutions on a hemacytometer to determine CFU/ml.

- Adjust the concentration of the stock with sterile PBST to yield $10^9$ CFU/ml for mice or $7 \times 10^7$ CFU/ml for guinea pigs, based on the needs of the specific inhalational chamber being used. See preparation notes, below.

- Conidial stock should be stored at 4°C until use. [Conidial stock may be used up to 24 hours after it was initially prepared. However, conidia should be recounted prior to infection to re-confirm concentration.]
• Confirm inoculum numbers by plating 100 µl on PDA plates overnight at 37°C using the following dilutions: $10^{-5}$, $10^{-6}$ and $10^{-7}$ for murine studies OR $10^{-4}$, $10^{-5}$ and $10^{-6}$ for guinea pig studies.

• **Preparation of final inoculum for particular chambers:**
  - Acrylic chamber:
    - Prepare a final inoculum consisting of at least 13 ml of $1 \times 10^9$ conidia per ml in PBST for mice and $7 \times 10^7$ conidia per ml for guinea pigs. Note that each run will require 12 ml for infection for a single 1 hour run of the chamber.
  - Madison chamber:
    - Prepare a final inoculum consisting of 20 ml of $1 \times 10^{10}$ conidia per ml in PBST. Note that each run will require approximately 12-15 ml for infection for a single 1 hour run of the chamber.

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**Strain Preparation Timeline**

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D-10 & D-7 & D0 & D+1 \\
Inoculate 1^{st} set of plates & Inoculate 2^{nd} set of plates & Day of infection Harvest Conidia & Overnight inoculum check \\
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7. Attachments
   - N/A

8. Deliverables
   - For the purposes of this SOP, two similar deliverables are possible. First, for the Acrylic chamber, is the preparation of a final inoculum consisting of at least 13 ml of $1 \times 10^9$ *Aspergillus fumigatus* conidia per ml in PBST for mice or a final inoculum of $7 \times 10^7$ conidia per ml for guinea pigs. Each run will require 12 ml for infection for a single 1 hour run of the acrylic chamber.

   The second choice, specific to the Madison chamber will be a final inoculum consisting of 20 ml of $1 \times 10^{10}$ conidia per ml in PBST. Each run will require approximately 12-15 ml for infection for a single 1 hour run of the Madison chamber.

9. References

   Vallor AC, Kirkpatrick WR, Najvar LK, Bocanegra RC, Kinney MC, Fothergill AW, Herrera ML, Wickes BL, Graybill JR, Patterson TF.

10. History
   Version 1.00. Original
   Version 1.10. Revisions made to text for purposes of clarification and uniformity.

11. Examples of Deliverables
    N/A