Spore suspension and Spore counting

Introduction

The purpose of this protocol is to obtain the spores generated from a plate culture in agar.

Materials

- MilliQ (MQ) water
- Mira-cloth
- Autoclaved Funnels
- Falcon tubes (50 mL)
- Cell spreader
- Optical Microscope

Procedure

Spore suspension

- 1. Pour MQ water into the fungal culture plates.
- 2. Rub the plate with the spreader. Make sure the spores get into solution. Spores are hydrophobic and can spread easily into the surroundings.
- 3. After spores are in suspension, if more plates with the same spores are to be harvested, pour water and spores to the next plate. Repeat steps 1-2 for as many plates as you have.
- 4. Set a filter (funnel + mira-cloth) on a Falcon tube.
- 5. Pour MQ water + spores onto the mira-cloth and filter solution. Make sure all liquid goes through the filter.
- 6. Do a second filtration on a new Falcon tube and filter. Make sure all liquid goes through.

Spore counting

- 1. Make 1:100 dilution with the spore suspension (only 10 μL needed).
- 2. With a counting chamber under the optical microscope, put 5 μL of the dilution into the centre of the chamber. Count spores in one of the squared cells.
- 3. Calculate spore concentration.

A. niger Glycerol stock

Introduction

The purpose of this protocol is to prepare a stock of *A.niger*.

Materials

- An agar plate with A. niger
- 50% glycerol
- Appropriate media
- Eppendorf tube or storage tube
- Cryo-tube
- MilliQ water

Procedure

- 1. Pour 2.5 mL of 50% glycerol on the plate.
- 2. Use a cell spreader to harvest spores.
- 3. Transfer 2 mL to an Eppendorf tube or storage tube (for the -20°C stock).
- 4. Pour 750 μ L of MilliQ water to a cryo-tube and transfer 500 μ L from the spore suspension (for the -80°C stock).