



STRAIN DEVELOPMENT

1. Before setting up a cross grow the strains for 2-3 days on YE media (or selection media if necessary).
2. Check the strains all the markers and phenotypes. This will ensure that you do not use the wrong strain.
3. Patch the strains on a

Calculating spore concentration using a haemocytometer.

1. Do a 1000X dilution of your spore sample. Load 10ul on the haemocytometer slide as follows.
2. To load the sample first place a coverslip in the haemocytometer slide. Then gently release the sample from the pipette tip into the wedge on the slide, making sure the coverslip is on top. Wait a few seconds for the sample to spread out properly.
3. Count the spores in each of the 4 (16 squared) corners of the slide.
4. From this calculate the average number of spores in a 16 squared corner.
5. The volume of the 16 squared corner is $1\text{mm} \times 1\text{mm} \times 0.1\text{mm}$. This is 0.1 c.mm or 0.1ul. Thus the spore concentration is average number of spores calculated above/0.1ul. Compute the actual spore concentration by taking into account the