

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 1mm X 1mm X 0.1mm. 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 initial dilution you manea4-5(v)10(e).the sample veye 537.07 Tm[T)-8(h)-3(i3 Tm9il)4(u)-3(tio) BD0