Microarray Technology
Lab Manual

Instruction Team:

Scott Tighe, Tim Hunter, Pat Reed, and Janet Murray

Manual: Version cerevisiae 12/09
Overview

In this module, you will learn how gene expression in yeast (*Saccharomyces cerevisiae*) is changed after exposure to oxidative stress induced by the addition of hydrogen peroxide (H$_2$O$_2$), an oxidizing agent. In the experiment, yeast are grown for 48 hours in 1/2x YPD broth containing 3x glucose. The culture is split into a control and treated group. The experiment culture is treated with 0.5mM H$_2$O$_2$ in Hanks Buffered Saline (HBSS) for 1 hour. The control culture is treated with HBSS only. Total RNA is extracted from both cultures and is converted to a labeled cDNA product through a multi-step process. The final synthesis product is taken back to the UVM Microarray Core Facility and hybridized to the Affymetrix yeast GeneChips. The resulting gene expression data are uploaded into bioinformatics data analysis software. You will select specific genes whose expressions have changed as a result of the exposure to H$_2$O$_2$ and generate a report to the class.
Why Use Yeast (*S. cerevisiae*)??

- easily manipulated in the laboratory
- simple eukaryote, unicellular
- rapid growth (doubling 1.5 - 2.5 hours)
- non-pathogenic
- stable haploid and diploid states
- complete genome sequenced
- very well characterized
- many pathways are conserved

Yeast Life cycle

http://www.bath.ac.uk/bio-sci/images/profiles/wheals2.gif
Special Notes- Important!

1] **RNase-free water** is DEPC water and the terms maybe used interchangeably throughout the protocol.

2] Record all data in notebooks.

3] Label the tops and sides of tubes with:
   
   - Sample ID/Initials
   - Date
   - What is in the tube…ie cDNA, RNA, etc.
   - Concentration

4] Check off each step in the lab manual as you complete it.

5] Read the Technical Discussion section before each day. This is fair game for quiz questions.

6] MSDS safety sheets are available for each chemical in the front of the room.

7] RPM on a Centrifuge does NOT equal G-force. See the conversion chart below

8] Use RNase Zap anytime you want, better safe then sorry!
Microarray Data Sheet

Individual identification:
Name____________________ (Group ID ______________________)
Experiment type________________________

1) Total RNA data from DAY 1
   RNA conc:___________ ug/ul:  A$_{260}$/A$_{280}$:__________

2) cDNA synthesis and clean-up:
   Volume RNA used: _________
   µg RNA used$^{[B]}$:__________
   Resuspension Volume ________
   Visible Pellet ________

3) RNA InVitro transcription:
   Volume of cDNA used in IVT reaction: ________

4) cRNA clean-up:
   Final volume$^{[C]}$: _________ ul
   Concentration: ___________ ug/ul
   Total RNA yield ___________ ug$^{[A]}$
   Final Adjusted RNA concentration: _________________ ug/ul

5) cRNA Fragmentation:
   Amount Fragmented: ____________
Attach Spectrophometer data

Attach gel images
Set-up before day 1: For course Instructor

Note: The broth culture must be inoculated about 48 hours before the treatment procedure of day 1 with a log phase culture of the yeast [See appendix A].

Note: The H2O2 treatment procedure must be performed 0.5 hour before class begins: total treatment time is 60 minutes.

Protocol for Preparing Yeast Cultures for Microarray Analysis  
(3-4 days prior to day 1)

Necessary Supplies
Autoclave (tape, tinfoil)  
Inoculating loop (or sterile swabs) and Bunsen burner  
Sterile flasks containing stir bars that are the same size: (1) 500 ml flask, (2) 125 ml flasks  
(2) Stir plates  
5 and 25 ml pipettes  
(sterile) Tape and sharpies for labeling

Supplies provided by VGN
Yeast Stain Saccharomyces cerevisiae [NRRLY-12632 or ATCC 18824]  
1/2xYPD+3x glucose nutrient media (modified powder –Fisher Scientific–This is pre-made by VGN staff. (12.5gram YPD powder+10g glucose+500ml water)  
30% Hydrogen Peroxide (H2O2)  
Sterile DI Water  
HBSS without Ca, Mg, or dye

The broth culture must be inoculated 48 hours before the treatment procedure of day 1. Incubation should be conducted in a room that is 70-80°F. It is best to inoculate this culture from a pre-culture that is in log phase.

Plan the treatment of the cultures so that the 1 hour treatment is complete 30 minutes AFTER the start of class on Day 1.

Check off each step in the protocol after completing it.

Please read through all instructions before proceeding.
A) Making the growth media and sterilizing flasks with stir bars.

___1) Prepare 250 ml of 1/2x YPD-3x Glu broth. Weigh 22.5 gm of the media and transfer to a 500 ml flask. Transfer 250 ml dd water to the flask. Place a stir bar in the flask, wrap tin foil loosely around the top, label the flask, and autoclave under standard conditions (15 psig at 121°C) for 15 minutes.

___2) Also autoclave (2) empty 125 ml flasks with the same size stir bar in each. Wrap top in foil.

It is important not to autoclave LONGER than 15 minutes as the sugars will caromatize and degrade.

B) Inoculating the parent culture

___1) Use sterile technique throughout this procedure. Bacterial contamination will be undetectable throughout the rest of the experiment and will adversely affect the final results.

___2) The parent culture should be inoculated 48 hrs before class of day 1 from a log phase pre-culture. See appendix A. Use a good size inoculum!!

___3) Transfer a colony of an actively growing yeast using an inoculating loop to the broth. Alternatively, if a liquid pre-culture is used, a 500 µl aliquot can be transferred to the new media.

___4) Uncap and tilt the 500 ml flask at a 30 degree angle [or so] and inoculate the 1/2x YPD-3x Glu broth.

Notice: you want to minimize the time the flask is open AND you do not want your hands or sleeves over the flask opening. Broth is easily contaminated and it will not be possible to detect this until the very end of the experiment.

___5) Flame the mouth of the flask and place the tinfoil cap back on the flask. Replace the tinfoil around the mouth of the flask, but not too tightly as it is necessary for oxygen to get in. It is important that the yeast grow aerobically as respiration is the metabolic process that builds cell mass.

___6) Place the flask on stir plate and stir approximately 48 hours at a medium speed at room temp (22-25°C). Do not have the speed so high that there is foaming. The culture should be aerated to ensure the yeast are not in a fermentation metabolism.
C) Treatment of yeast cultures (must be started 0.5 hr before class on day 1)

**Preparing working solution of H_{2}O_{2} to achieve a final concentration of 5.0 mM in culture flask**

**Preparation of H_{2}O_{2} Stock Solution:** Combine 5.0µL of 30% H_{2}O_{2} and 495 µL of Hanks Balanced Saline Salt [HB without phenol red, Mg, or Ca]. Vortex to mix well.

**Working H_{2}O_{2} solution:** Make a 1:10 dilution by combining 100µL H_{2}O_{2} stock solution with 900µL HBSS and vortex. Use this for spectrophotometer measurement and for the experiment.

Measure the absorbance at 240nm.

Blank the spectrophotometer with HBSS and determine the absorbance of the working solution.

\[ A_{240} = \text{___________} \]

**Concentration of H_{2}O_{2} in Working Solution:**

\[ (A_{240}) \times 229 = \text{_____ mM of H}_{2}\text{O}_{2} \text{ in working solution} = A \]

Volume of working H_{2}O_{2} solution to use in experiment to achieve a 5.0 mM final concentration in a 25 ml culture volume.

\[ \text{ul of working solution to use} = \frac{5.0 \text{ mM}}{A \text{ mM}} \times 25\text{ml} \times 1000\text{ul/ml} \]

where A = concentration of H_{2}O_{2} in working solution as determined above.
1) From the 48 hour old culture, aseptically transfer 25 ml of the broth culture into two (2) sterile 125 ml flasks each containing a stir bar using a sterile 50 ml pipet.

2) Label one flask as “treated” and aseptically add enough H₂O₂ from the “working stock” to achieve a final H₂O₂ of 5.0 mM [See below]. Label the control flask as “control” and add an equal amount of HBSS as you did H₂O₂.

3) Place each flask on separate stir plates and stir at room temperature for 1 hour at [as close as] the same speed as possible. Cultures should be ready for harvest 30 minutes after the start of the class period. Therefore, H₂O₂ is added 30 minutes before the start of the class.
DAY 1

Isolating Total RNA from the Yeast Using Enzymatic Lysis Protocol.

Technical Overview:

You will be treating the yeast, *Saccharomyces cerevisiae*, with hydrogen peroxide \([H_2O_2]\) to determine the effects on gene expression during oxidative stress conditions. After one hour of treatment the treated and control yeast cultures are pelleted by centrifugation. The yeast are treated with lyticase, an enzyme that degrades the poly-\(\beta\)-1,3 glucan bonds in the cell wall, resulting in the formation of spheroplasts (a yeast cell without a cell wall). The second step ruptures the spheroplasts using a buffer containing the chaotropic agent, guanidinium isothiocyanate, which dissolves the cell membrane. All the RNA is collected using a silica gel spin column manufactured by Qiagen. (RNeasy system)

Please read through all laboratory procedures prior to lab.

Check off each step in the protocol as you complete it.
The Silica Membrane Chemistry

Guanidium isothiocyanate (GITC) is a chaotropic salt at about 5 M in the RLT buffer. This solution (pH 6.4) used in conjunction with ethanol and water disrupts the molecular characteristics of water and creates conditions where RNA is less soluble. This promotes nucleic acids to electrostatically bind to the silica membrane of the spin column. A stoichiometric balance of GITC, ethanol, pH, and water controls the binding of either RNA or DNA to the silica. Therefore, changes in this stoichiometry will select a majority of either one or the other. Since this not a 100% selective procedure DNase treatment to destroy any DNA is required. The use of ethanol in rinse RPE buffer maintains the bond between the silica and RNA. RNA is eluted from the membrane with pure water, which removes the salts and breaks the electrostatic bond of RNA with the silica. RNA in an almost pure form is collected by centrifugation.

![Diagram of the binding of RNA to the silica membrane](image.png)

**The RNA is electrostatically bound to the silanol groups of the silica membrane using an isothiocyanate-guanidium salt bridge to the negatively charged phosphate backbone of the RNA.**
Isolating total RNA from *Saccharomyces cerevisiae* using Enzymatic Lysis

**Materials:**
- Axygen MCT150C 1.7ml tubes
- P1000, P200, and P20 pipets
- Yeast Lytic Enzyme-VWR # IC15352710-MP Biomedical 153527-10,000 units
- 100% Ethanol
- DEPC water (aka RNase-free water)
- Qiagen Mini RNeasy kit
- Qiagen DNase Kit
- Vortex
- Microcentrifuge
- Spectrophotometer
- Eppendorf UVettes
- 2 ml microcentrifuge tubes
- 2% SDS

1. Prepare an RNase-free work zone using RNase ZAP. This will be demonstrated by the instructor.

   **Note: The Instructor will perform step 2 tasks.**

2a. Prepare fresh working lyticase reagent by adding 1 ml of DEPC water directly to the lyticase vial. Vortex well and invert several times to insure complete mixing. This 10 units/µl solution is stable for 12 hours.

   10,000 U Lyticase
   1 ml DEPC Water

2b. Prepare fresh DNase I solution using the Qiagen DNase kit and store on ice.

   10 µl DNase I
   70 µl Buffer RDD
   80 µl per reaction

3. Label a 1.7 ml microcentrifuge tube with your identification information. Transfer 1.5 ml of the appropriate yeast culture the tube.
4. Centrifuge the tube at 5000x g (about 3/4 full speed) for 2 minutes at room temperature. Carefully remove the supernatant (without disturbing pellet) using a micropipette and discard the supernatant.

**** Repeat step 4 if pellet size is too small or if indicated by instructor.****

5. Add 1000 ul DEPC water to the pellet, vortex, and centrifuge at 5000 x g for 2 minutes. Discard the supernatant. Remove as much of the liquid as possible from the yeast pellet.

6. Add the following to the yeast pellet and vortex to mix.

<table>
<thead>
<tr>
<th>Buffer</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>SG buffer</td>
<td>10 µl</td>
</tr>
<tr>
<td>Lyticase solution</td>
<td>30 µl</td>
</tr>
</tbody>
</table>

7. Incubate for 30 min at room temperature.

8. Gently swirl the tube every 10 minutes to generate spheroplasts. Spheroplasts must be handled gently.

[optional] …Check spheroplast formation by placing 10ul of cells on a slide with 10ul of 2% SDS. These can be swirl mixed with a pipet tip right on the slide.

9. Add 350 µl b-RLT buffer to the tube and vortex vigorously for 1 minute. Ensure your tube is tightly capped by holding the lid closed while vortexing. This procedure will lyse the spheroplasts.

10. Add 250 µl 100% ethanol to the tube and briefly vortex. A precipitate may form after the addition of ethanol, but this will not affect the collection of RNA.

11. **Collecting the RNA:** Assemble an RNeasy spin column with a 2 ml collection tube. Label the spin column with your identification information. Carefully transfer all of the solution from step 10 to the spin column using a micropipette. Be careful not to touch the silica membrane with the pipette tip. Close the tube and centrifuge for 15 seconds at full speed. The RNA in the sample will adhere to the silica membrane of the spin column.
12. Remove the spin column from the collection tube. Pipet the pass through liquid (the liquid in the collection tube) back onto the spin column and centrifuge again. Discard the collection tube (with the flow through fluid) and place the spin column into a new 2 ml collection tube.

13. Add 350 µl RW1 buffer to the spin column. This solution is used to wash the RNA and remove salts and dissolved cellular debris. Close the tube and centrifuge for 15 seconds at full speed.

14. **Degrading DNA:** Remove the spin column from the centrifuge, open the tube lid and add 80 µl of DNase I solution to the middle of the spin column membrane. This step will degrade any DNA in the sample and dissociate it from the silica membrane. Close the column lid and incubate at room temperature for 15 minutes.

15. Transfer the spin column to a new 2 ml collection tube.

16. **Cleaning and washing the RNA:** Add 350 µl RW1 buffer to the spin column and centrifuge at full speed for 15 seconds.

17. Discard the collection tube and place the spin column into a new 2 ml collection tube.

18. Add 500 µl RPE buffer to the spin column to wash the RNA on the silica membrane. Close the tube and centrifuge for 15 seconds at full speed.

19. Discard the collection tube and place the spin column into a new 2 ml collection tube.

20. Wash the silica membrane again by adding another 500 µl RPE buffer to the spin column. Close the tube and centrifuge for 15 seconds at full speed.

21. Place the spin column in a new 2 ml collection tube and centrifuge in a microcentrifuge at full speed for 1 minute to ensure any residual liquid is removed from the silica membrane.

22. **Recovering the RNA:** Label a 1.7 microcentrifuge tube with your identification information. Transfer the spin column to this tube. The lid of the microcentrifuge tube will be off to the side. Carefully pipet 30 µl DEPC water directly onto the very center of the silica membrane. **Do not touch the silica membrane with the pipet tip (see diagram).** Look closely as you perform this step. Use both hands when pipetting, one on top and one down by the tip to guide the pipet. Make sure the water is evenly distributed on the membrane.
23. Incubate the spin column at room temperature for 1 minute. Centrifuge at full speed for 30 seconds.

24. Carefully transfer the 30 µl that is recovered in the 1.7 ml tube back onto the center of the silica membrane of the same spin column as in step 22 above. Place the column back in the same 1.7 ml collection tube and incubate for 1 minute at room temperature. Centrifuge for 1 minute at full speed. This double extraction ensures that the maximal amount of RNA is eluted from the membrane.

25. Label two new 1.7 microcentrifuge tubes with your identification information. Transfer all the recovered RNA sample to one of these tubes.

26. Transfer 4 µl of this sample to the other labeled tube. This sample will be transported back to UVM for Nanodrop quantification and RNA assessment.

27. Place all samples on ice.

28. Quantify the RNA: Measure the absorbance of the RNA sample on the spectrophotometer at a 1 to 50 dilution (1 µl sample + 49 µl H2O). This will be demonstrated by the instructor.

Absorbance of sample: ____________ A260/A280 ratio: ________________
Concentration of RNA in sample: ________________ ug/ul

Record these data on the data sheet provided on page 5 of the manual.

29. Evaluate the RNA quality using agarose gel electrophoresis. This will be done as a class.

30. Set up the E-gel electrophoresis apparatus. Pre-run the gel for 2 minutes by pressing the button continuously for 10 seconds. You will see the light start to blink indicating it is in a pre-run mode. It will beep and stop after 2 minutes.

Add 14 µl of water to each well on the E-gel. Add 1 µl of each sample to each well and pipet up and down to mix well. Record which sample is in each lane. Run the gel for 20 minutes.

30. Take a photo of the gel.

A sample of your RNA may be analyzed on the Agilent 2100 Bioanalyzer at UVM, which provides a more sensitive image of the RNA quality. (Described on the next page).
**RNA Analysis using the Agilent 2100 Bioanalyzer**

The Agilent bioanalyzer 2100 is an automated instrument that employs the use of Lab-on-a-Chip technology for analyzing RNA. This provides highly resolved data regarding the integrity of the RNA. The system is similar to an agarose gel where RNA is added to a well containing a gel and high-resolution electrophoresis is conducted through microfluidic channels. The use of an intercalating dye allows the RNA to be detected with an Nd-YAG diode-pumped laser.

![Image of a computer and Agilent bioanalyzer](image)

**Moderately Degraded RNA**

Elevated and "rough" baseline [shiny]. Significantly reduced 25S ribosomal peak indicating degradation.

**Fully Intact RNA**

Clean flat baseline and tall sharp peaks for 18S and 25S ribosomal RNA. 25S equal to or higher than that of 18S.
Day 2

Microarray Target Preparation- cDNA synthesis

Technical Overview:

Today you will be starting to prepare the target cRNA for the Genechip analysis. This is a lengthy procedure and will be broken down over several lab periods. You will be starting with mRNA, which comprises 0.5-2% of the total RNA in typical eukaryotic cells. First, you will generate the first complementary DNA strand (cDNA) from the mRNA. This is called the first strand cDNA synthesis. The mRNA molecule is oriented in a 5’ to 3’ direction. This step utilizes a primer with a sequence of 24 T's in a row and is called oligo d(T). This primer binds to the poly A tail of mRNA (the 3’ end). The primer also contains the consensus-binding site for T7 RNA polymerase (T7 promoter sequence) which is 36 bases long. The total length of the primer is 60 bases. Once the primer has hybridized to the poly-A tail of the mRNA molecule, reverse transcriptase, an enzyme that uses RNA as a template to synthesis DNA, is added and the first strand of cDNA is synthesized. The final product of this reaction is a hybrid molecule composed of one RNA strand (the original mRNA molecule) and one DNA strand. The first DNA strand is synthesized in a 3’ to 5’ orientation relative to the mRNA strand (see the diagram on page 20, step 2). It is important to note that rRNA and tRNA molecules in the sample do not have a poly-A tail and will not bind the oligo T primer and will not be used as a template for cDNA synthesis in this procedure.

The full sequence of T7 Oligo d(T)24 is:

\[
5'\text{-GCCGAGTGAAATTGATACGACTCCTATAGGAGCCGG-TTTTTTTTTTTTTTTTTTTTTTTTTTTTT - 3'}
\]

\(d(T)_{24}\)-Binds to Poly-A of mRNA
The next step in the cDNA reaction is the synthesis of the second cDNA strand from the first cDNA strand. This is done by adding RNase H, DNA polymerase I, and DNA ligase. The RNase H causes many nicks in the phosphodiester bonds (between the phosphate and the ribose sugar) of the bound mRNA strand (not the cDNA stand already made) creating exposed 3’OH groups. DNA polymerase I enzyme uses these 3’ OH groups as a priming site; displacing the RNA in the strand and creating another cDNA strand using the first cDNA strand as a template. It continues in this fashion until it reaches another nicked site, where it stops. The DNA polymerase I can only displace RNA, not DNA. After cDNA has been synthesized from all the nicks, the 3’OH nicks are reconnected or ligated. This is accomplished using DNA ligase enzyme. After the two hour synthesis process, all the RNA has been replaced with cDNA and ligated to form one double stranded cDNA molecule. However, because the DNA polymerase does not synthesize completely to the end of the strand, a T4 DNA polymerase is added. This enzyme ensures the full synthesis of the T7 promoter site from the original T7 oligo d(T) primer used in the first synthesis reaction. The final product is a double stranded cDNA molecule which includes the original mRNA sequence (in DNA form) on the second strand plus the full T7 RNA polymerase binding site at the 3’ end of the molecule—the dark bar shown on step 3 on page 20.
Synthesis of cDNA:

The starting material for your target preparation will be high quality, fully intact, total RNA from yeast you harvested on Day1. The minimum concentration to start with is 100-270 ng/µl. After isolation and purification of the yeast RNA, the concentration is determined by absorbance at 260 nm ($A_{260}$) on a spectrophotometer (1.0 A = 40 ng/ul RNA). The $A_{260}/A_{280}$ ratio should be approximately 1.8-2.1. We are checking the quality of the RNA by running an aliquot on the Agilent Bioanalyzer 2100 (performed prior to Day 2 class at UVM) and running on an agarose gel (EGEL) prior to starting the assay.

Overview of cDNA Synthesis Reactions
Day 2

Microarray Target Preparation- cDNA synthesis

First Strand cDNA Synthesis

Note: You will be measuring and transferring very small volumes. Centrifuge all reagent tubes before starting as some have only 1 µl in them. When pipetting, look closely at your pipet tip to ensure you have picked up the sample and look at the tube to which you are delivering it to ensure it has been added.

_____ 1. Label a 0.5 ml PCR tube with your identification information. From the concentration of your RNA determined on Day 1, calculate the volume of the sample needed to contain 3.0 ug. Transfer this amount to the tube. Add DEPC water to bring the volume to 11.0 ul.

Example: RNA concentration of your sample = 800 ng/ul or .8ug/ul.
3.0 ug/.8 ug/ul = 3.75 ul. Transfer this amount to the tube.
11.0 ul – 3.75 ul = 7.25 ul = volume of DEPC water to add to tube.

- RNA concentration of your sample: _____________ ug/ul
- 3.0 / (your RNA concentration) = _____________ ul. This is the volume of your sample you will add to the tube.
- 11.0 – (the volume of your RNA sample) = ___________ ul. This is the volume of DEPC water to add to the tube.

_____ 2. Add 1.0 ul T7 oligo d(T)24 reagent to the tube. Make sure you see the reagent transfer. Vortex the tube and centrifuge at full speed for 5 seconds. Place the tube in a thermocycler set at 70°C for 10 minutes.

_____ 3. While the 70°C incubation is in progress prepare the following master mix in a new tube. Add the following reagents IN ORDER to make the first strand master mix. Vortex and centrifuge at full speed for 5 second and place the tube on ice.
First strand master mix

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>First Strand Buffer 5X</td>
<td>4 µl</td>
</tr>
<tr>
<td>0.1M DTT</td>
<td>2 µl</td>
</tr>
<tr>
<td>10mM dNTP</td>
<td>1 µl</td>
</tr>
<tr>
<td>Superscript II</td>
<td>1 µl</td>
</tr>
</tbody>
</table>

Use a P2 micropipette to measure volumes of 2 ul or less.
4. After the 70°C incubation in step 2 is finished, add the first strand master mix made in step 3 to the tube and incubate in a thermocycler at 42°C for 60 minutes. When first strand synthesis incubation is finished, place the tube on ice. During this incubation, prepare the second strand master mix.

**Second Strand cDNA Synthesis**

1. Make the following master mix in a new tube. Keep it on ice. Centrifuge all reagent tubes before using to ensure the contents are at the bottom of the tube. Add the reagents in the order listed.

   Second strand master mix
   
<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEPC Water</td>
<td>91 µl</td>
</tr>
<tr>
<td>5x Second Strand Buffer</td>
<td>30 µl</td>
</tr>
<tr>
<td>dNTP(10mM)</td>
<td>3 µl</td>
</tr>
<tr>
<td><em>E.coli</em> DNA Ligase (10U/µl)</td>
<td>1 µl</td>
</tr>
<tr>
<td><em>E.coli</em> DNA Polymerase I (10U/µl)</td>
<td>4 µl</td>
</tr>
<tr>
<td><em>E.coli</em> RNase H (2U/µl)</td>
<td>1 µl</td>
</tr>
<tr>
<td>Total volume to be added to First Strand tube</td>
<td>130 µl</td>
</tr>
</tbody>
</table>

2. Vortex the tube and centrifuge for 5 seconds at full speed.

3. When the 42°C incubation is finished, transfer all 130 µl of the second strand master mix to the first strand synthesis tube. Vortex and centrifuge for 5 seconds at room temperature. Incubate the tube for 2 hours at 16°C in a thermocycler.

The next two steps may be performed by the instructor

4. At the end of the 2 hour incubation and while the sample is still at 16°C, add 2µl T4 DNA polymerase reagent to the tube. Mix the tube well by inversion. Incubate at 16°C for 5 minutes. Do not incubate longer then 5 minutes or the quality of the cDNA may decrease due to the 3’ to 5’ exonuclease activity of the polymerase.

5. At the end of the 5 minute incubation, add 10 µl 0.5 M EDTA to the tube to stop the T4 DNA polymerase reaction. Store the tube at -20°C.
Day 3

**Technical Overview: Cleaning the cDNA:**

The double stranded cDNA generated on Day 2 must be cleaned and purified before proceeding to the next step. This is done using a solution of phenol-chloroform-isoamyl alcohol (PCI) at a alkaline pH to denature proteins and trap them in the organic phase of the reagent. cDNA remains in the aqueous phase and the two phases are separated using a phase lock gel system. The phase lock gel tube is centrifuged causing the gel in the tube to migrate between the aqueous phase (on the top and containing the cDNA) and the organic phase (on the bottom and containing the denatured proteins). Recovery of cDNA is easy using this method. The pH of the PCI solution is maintained by Tris buffer at pH 8.3 to ensure the cDNA will dissolve in the aqueous layer.

cDNA is purified from the aqueous phase through a precipitation step using ethanol [an alcohol] and ammonium acetate [a salt]. Because DNA is negatively charged, a high salt concentration is used to mask the charge, decreasing the solubility of the cDNA. Ethanol is a non-polar solvent and DNA will not readily dissolve in it in these conditions and precipitates from the solution. This reaction is sometimes done at -20°C to encourage maximum DNA precipitation. After precipitation, the cDNA is centrifuged at high speed to form a pellet with a visualization agent called Pellet Paint, an additive that contains glycogen (a co-precipitate) and a dye. This helps generate a high quality visible pellet on the bottom of the tube. The cDNA pellet is washed several times with 80% ethanol, dried, and resuspended in DEPC water.
The In vitro Transcription (IVT) Reaction:

The IVT step uses the T7 promoter site that is attached to the T7 Oligo d(T)24 primer used in the first strand cDNA synthesis. T7 RNA polymerase enzyme binds to this site and generates cRNA molecules. In this arrangement the original mRNA sequence is the template strand and the cRNA generated is antisense relative to this sequence. Two unique nucleotides are used in the reaction. A select proportion of the C and U ribonucleotides have molecules of biotin attached. A and G nucleotides are not biotinyated. As a result, the cRNA generated has random amounts of biotin label incorporated into its structure. The biotin will be the beginning of the signal generation process used on the gene chips based on a strepavidin-phycoerytherin reaction. During the IVT reaction the T7RNA polymerase binds to the T7 promoter site and generates a molecule of cRNA. During the 16 hour incubation time for the IVT, a single enzyme can generate many thousand cRNA molecules from the cDNA template. The net result is a total increase in RNA at the end of the reaction. For a typical IVT reaction, 1.5 to 2.5 ug/ul yields are expected. 10 ug of cRNA is carried forward to Day 4 for fragmentation and hybridization.
Safety Considerations for Phenol Chloroform Isoamyl:

Note: Phenol Chloroform Isoamyl Alcohol is at a ratio of 25:24:1 and saturated with 10 mM Tris- HCl pH 8.3. The Tris layer floats on top of the PCI layer. Phenol is an organic acid and care must be used when handling. Phenol can burn skin and be absorbed into the circulatory system and cause health problems. Chloroform is an organic solvent and should be used in a hood.

- **Phenol**
  - Corrosive
  - Local anesthetic
  - Can severely damage eyes and skin
  - Can cause severe burns of the skin and eye
  - Poisonous if ingested—can be fatal

- **Chloroform**
  - Can cause irreversible corneal injury
  - Fatal if swallowed
  - Flammable

- **Isoamyl Alcohol**
  - Flammable
  - Poisonous

Note: PCI tends to leak out of the pipet tips so work quickly when measuring it or transferring it from one container to another.
Day 3

Precipitating the cDNA:

1. The total volume of the cDNA synthesis step from Day 2 is 162 µl. The PCI step requires equal volumes of aqueous and organic mixtures. Centrifuge a Phase Lock Gel tube at full speed for one minute to insure the gel is at the bottom of the tube. DO NOT VORTEX PHASE LOCK TUBES.

2. Add 162 µl of the bottom layer from the Phenol Chloroform Isoamyl Alcohol (PCI) to the contents of the cDNA synthesis reaction tube from Day 2 and vortex for 5 seconds to mix the contents.

   Note: PCI tends to leak out of the pipet so work quickly and hold the cap tightly as leaking can occur during the vortex step.

3. Transfer all of the cDNA-PCI mixture to the phase lock gel tube using a micropipet. DO NOT VORTEX the phase lock gel tube.

4. Centrifuge at full speed for 2 minutes.

5. Label a new 1.7 microcentrifuge tube. Use a micropipette to transfer the top layer from the phase lock gel tube to the tube. Try to collect as much of the layer as possible.

6. Add the following to the 1.7 ml microcentrifuge tube and vortex.

   - Ethanol (100%) 405 µl
   - NH₄OAc (7.5M) 80 µl
   - Pellet Paint 1 µl

7. Place the tube in the centrifuge with the hinge of the tube facing out and centrifuge at full speed for 20 minutes at room temperature.

8. GENTLY remove the tube from the centrifuge being careful not to disturb the cDNA pellet. The pellet should be pink and approximately the size of a grain of salt and on the side of the tube under the hinge. Place on ice and immediately proceed to next step.
Is there a pellet?__________

**At this point a tiny pink visible pellet should be present on the bottom of the tube. If not, see the instructor.**

Cleaning the cDNA Pellet:

1. Using a P1000 micropipet, carefully remove the liquid from the tube. Tip the tube to remove as much liquid as possible, but don’t lose the pellet. Remember that the pellet is your sample!

2. Add 500µl ice cold 80% ethanol (store in -20°C freezer) to the tube. Gently cap the tube and invert it slowly several times. Watch your pellet very closely. Place the tube back in rack and let the pellet settle to the bottom of the tube for a minute. Alternately, you may centrifuge the tube at full speed for 15 seconds to get the pellet back down to the bottom of the tube.

3. Using a P1000 micropipet, carefully remove the ethanol being very careful not to disturb the pellet. Tip the tube to enable removal of as much liquid as possible.

4. Repeat steps 2 and 3 with a new aliquot of 80% ethanol.

5. Finally, remove all of the ethanol. To do this, remove as much as possible using a P1000 micropipet. Centrifuge the tube at full speed for 5 seconds and use a P20 micropipet to remove the last few microliters of liquid. The goal is to remove as much ethanol as possible without disturbing the pellet.

6. Place the tube on its side with the cap open in a drying box for 10 minutes to evaporate the remaining ethanol. The dried pellet is easily lost once it is dry. Be very careful to handle the tube gently. When done, close the cap gently. Visualize the dried pellet to confirm it is present in the tube.
7. Resuspend the pellet in 22 µl of DEPC water and place the tube on ice.

**In Vitro Transcription (IVT): Synthesis of biotin labeled cRNA from cDNA using the Enzo Kit**

1. The ENZO bioarray kit contains all the reagents needed to prepare biotin labeled cRNA from cDNA.

A master mix for the entire class will be prepared as follows. The instructor will prepare this mix or designate someone from the class. This must be done in an RNase-free area free from air currents.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Amt/sample</th>
<th>#Samples</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reagent 1 [10x Reaction buffer]</td>
<td>4 µl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reagent 2 [10x Biotin nucleotides]</td>
<td>4 µl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reagent 3 [10x DTT]</td>
<td>4 µl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reagent 4 [10x RNase Inhibitor]</td>
<td>4 µl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reagent 5 [20x T7 RNA polymerase]</td>
<td>2 µl</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Total Volume</strong></td>
<td><strong>18 µl</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**NOTE:** Prepare sufficient master mix for the number of samples plus one. This will ensure enough for pipetting purposes.

2. Label a 0.5 ml PCR tube. Combine the following in the tube and pipet up and down several times to mix. Spin in a centrifuge at full speed for 5 seconds to get all reagents to the bottom of the tube.

- cDNA mixture: 22 µl
- Enzo master mix [from above]: 18 µl
- Total Volume: 40 µl

3. Incubate the tube at 37°C for 16 hours in the thermocycler. When the reaction is complete, store the sample at -20°C.
Day 4

**Cleaning and fragmenting the biotinylated cRNA**

The biotinylated cRNA generated on day 3 must be cleaned and purified from the enzymes and reagents used in the IVT reaction. Removing the unused biotinylated NTPs, buffers and proteins is necessary as they interfere with quantitative measures. Proteins and NTP’s absorb at 260 and/or 280nm and will affect the quantitation of the cRNA. They also interfere with hybridization of the cRNA targets on the gene chip.

You will use an RNeasy spin column to clean the IVT product similar to that used on Day 1. The cRNA is washed and eluted from the spin column with DEPC water. The cRNA is analyzed with a spectrophotometer and the concentration and purity determined. An $A_{260}/A_{280}$ ratio of 1.8-2.1 indicates a clean cRNA sample that can be used for subsequent steps.

The cleaned cRNA is too large to hybridize onto the gene chip. It is fragmented into randomly short segments using a non-enzymatic, heat catalyzed magnesium salt reaction. The final product is assessed on an agarose gel.

**cRNA Fragmentation:**

\[10\mu g \text{ adj cRNA + fragmentation buffer} \xrightarrow{94^\circ C, 30'} \text{fragmented cRNA}\]

Run fragmented cRNA on precast E-Gel to assess fragmentation procedure and cRNA quality. The quality of cRNA is essential to obtaining valid results.
Day 4

Cleaning the biotinylated cRNA:

1. Remove the Day 3 tubes from the freezer and let thaw. Transfer the entire cRNA sample to a new 1.7 ml microcentrifuge tube. Add 60 µl of DEPC water and 350 µl BME-RLT and vortex for 5 seconds.

2. Add 250 µl 100% ethanol and vortex again.

3. Label an RNeasy spin column and transfer the entire cRNA sample to the column. As before, be careful not to touch the tip of the pipet to the silica membrane. Place the spin column in a 2 ml collection tube, close the lid and centrifuge for 15 seconds at full speed.

4. Remove the spin column from the collection tube. Pipet the pass through liquid (the liquid in the collection tube) back onto the spin column and centrifuge again for 15 seconds. Discard the collection tube (with the flow through fluid) and place the spin column into a new 2 ml collection tube.

5. Pipette 500 µl of RPE buffer onto the spin column. Close the tube and centrifuge for 15 seconds at full speed. Discard collection tube and the pass through liquid. Place the spin column into a new collection tube.

6. Add another 500 µl RPE buffer to the spin column. Close the tube and centrifuge for 15 seconds at full speed. Discard the collection tube and pass through liquid.

7. Transfer the spin column into a new collection tube and at full speed for 1 minute.

8. Label a new 1.7 ml microcentrifuge tube with your identification information. Transfer the spin column to this tube.

9. Pipet 30 µl DEPC water onto the RNeasy silica-gel membrane and wait 1 minute. Make sure the DEPC water contacts the entire membrane. Be careful not to touch the membrane with the pipet tip. Close the tube and centrifuge for 1 minute, at full speed.

10. Carefully transfer the 30 µl that is recovered in the 1.7 ml tube back onto the center of the silica membrane of the same spin column. Place the column back in to the same 1.7 ml collection tube and incubate for 1 minute at room temperature. Centrifuge for 1 minute at full speed. This double extraction ensures that the maximal amount of cRNA is eluted from the membrane.
11. Label a new 1.7 microcentrifuge tube with your identifying information. The old tube lid may have been contaminated in the last few procedures. Transfer the cRNA this tube.

12. Determine the cRNA concentration as on Day 1. Prepare a sample for analysis by adding 2 ul of your sample to 48 ul DEPC water. Determine the concentration and the $A_{260}/A_{280}$ ratio.

Absorbance $A_{260}$ of sample: ___________________

$A_{260}/A_{280}$ ratio: ___________________

Concentration of RNA in sample: ____________ ug/ul

Total yield of RNA = 30 ul x concentration = ________________ ug.

Record these data on the page 5 data sheet.
Calculating the Adjusted cRNA Concentration:

The final concentration of cRNA must be adjusted because the original amount of RNA you started with on Day 1 has been carried along in the experiment. Over 98 percent of that RNA was not mRNA but rRNA and tRNA. The amount of RNA measured above also includes these RNAs in the sample and must be adjusted to remove them from the final cRNA concentration.

13. Calculate the adjusted cRNA concentration using the following formula:

\[
\text{adjusted cRNA concentration = } \frac{[\text{AxB}] - \text{C}}{\text{B}}
\]

Calculation example:

- The amount of RNA started with on Day 1 = 3 ug (C in the formula)
- Concentration of RNA from step 12 above = 1.8 ug/ul (A in the formula)
- Volume of elute = 30 ul (B in the formula)

Total yield from step 12 above = 1.8 ug/ul x 30 ul = 54.0 ug (A x B in the formula)

Example math:

- The adjusted cRNA concentration = (54.0 ug – 3 ug)/ 30 ul = 1.7 ug/ul

The cRNA concentration in the sample is 1.7 ug/ul and this value is used in future calculations. Record these data on the page 5 data sheet.
**Fragmenting the cRNA for Target Preparation**

The cRNA molecules generated in the IVT reaction are too large to hybridize on the gene chip. In this step they are fragmented into randomly sized pieces of 35 to 250 bases in length. The cRNA is fragmented using a metal-induced hydrolysis reaction, referred to as a hammerhead cleavage. Both the unfragmented and fragmented cRNA are visualized on an agarose gel. This is important because you need to ensure the unfragmented cRNA has not degraded during the handling steps leading up to this point.

Affymetrix recommends the cRNA used in the fragmentation procedure be sufficiently concentrated to maintain a small volume during the procedure. This will minimize the amount of magnesium in the final hybridization cocktail. The cRNA must be at a minimum concentration of 0.6ug/ul. The fragmentation buffer contains a basic solution of TRIS acetate buffer, magnesium acetate, and potassium acetate, which causes a metal induced RNA hydrolysis at high temperature. See the diagram on the next page.

1. You will fragment 10.0ug of cRNA that will be used for gene chip analysis. To determine the volume of your sample needed in this step, divide 10.0 by the concentration of your adjusted cRNA sample.

Example: your adjusted cRNA concentration = 1.7ug/ul.

\[
10 \text{ ug} / 1.7 \text{ ug/ul} = 5.9
\]

Use 5.9 ul of your sample in the next step. (Record this value on the page 5 data sheet.)

The fragmentation reaction requires 16 ul of volume for the sample. The additional volume is made up with DEPC water. To determine how much DEPC water to use, subtract the volume of cRNA used from 16.

\[
16.0 \text{ ul} – 5.9 \text{ ul} = 10.1 \text{ ul of DEPC water used.}
\]

cRNA volume = ____________________ ul
DEPC water volume = ____________________ ul
2. Label a 0.5 ml PCR tube with your identification information. Transfer the amount of cRNA determined in step 1 above to the tube. (In the example this is 5.9 ul). Add DEPC water to bring the volume to 16.0 ul. (This is 10.1 ul in the example). Add 4.0 ul 5x fragmentation buffer to the tube. The total volume in the tube should be 20.0 ul.

3. Vortex the tube and centrifuge for 10 seconds. Incubate the tube at 94°C for 30 minutes in a thermocycler. Put on ice following the incubation.

**Fragmentation: Hammerhead Cleavage**

The intermediate product is a 2', 3' cyclic nucleotide. The next step is hydrolysis at the 3' carbon. The cyclic nucleotide is now a 2'-nucleotide. Hydrolysis at the 2' carbon produces the 3'-nucleotide.
**Agarose gel assessment of unfragmented and fragments cRNA samples:**

Use an agarose gel to assess the quality of your unfragmented and fragmented cRNA.

1. Set up the E-gel electrophoresis apparatus. Pre-run the gel for 2 minutes by pressing the button continuously for 10 seconds. You will see the light start to blink indicating it is in a pre-run mode. It will beep and stop after 2 minutes.

2. Add 14 µl of water to each well on the E-gel.

3. Add 2 µl of each sample to each well and pipet up and down to mix well. Run the fragmented and unfragmented cRNA of each sample next to each other. Record the identity of each sample in each lane.

4. Add 4 ul of a DNA ladder (size standards) to one lane of the gel

5. Run the gel for 20 minutes.


**Note:** The fragmented sample should be a smeared band at the bottom of the gel indicating fragments of RNA between 35 and 250bp. The unfragmented sample should be a smear from the top to the bottom of the gel as the cRNA molecules are of various lengths.

![Image of agarose gel](image-url)
To Be Performed at the University of Vermont

These steps will be performed at the University of Vermont Microarray Core Lab and will be shown as a powerpoint presentation during your class visit.

Preparing the Hybridization Mix

The hybridization mixture contains several internal controls that provide information about the success of the hybridization procedure. These controls are the B2 oligo for which there are several hundred probes along the outer edge of all expression arrays and a checkerboard pattern in each corner. These predefined patterns provide signals for the Affymetrix® Microarray Suite software to perform automatic grid alignment during image analysis. They can also be used to align the grid manually. The fluorescence intensities for control oligo B2 are not used for analyzing data. The bioB, bioC, and bioD are biotinylated gene fragments much like your sample which represent the biotin synthesis pathway from the bacteria *E. coli*. The cre control is the recombinase gene from P1 bacteriophage. These biological controls are prepared at standardized concentrations and signal intensities obtained on these genes provide information on how well the hybridization, washing, and staining procedures have performed.

1. Prepare the following:

<table>
<thead>
<tr>
<th>Component</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fragmented cRNA</td>
<td>[10µg]</td>
</tr>
<tr>
<td>Control B2 Oligo</td>
<td>3.3 µl</td>
</tr>
<tr>
<td>20x Eukaryotic Control mix [bio B, bio C, bio D, Cre]</td>
<td>10 µl</td>
</tr>
<tr>
<td>Herring Sperm DNA [10mg/ml]</td>
<td>2 µl</td>
</tr>
<tr>
<td>Acetylated BSA [50mg/ml]</td>
<td>2 µl</td>
</tr>
<tr>
<td>2x Hybridization Buffer</td>
<td>100 µl</td>
</tr>
<tr>
<td>Water</td>
<td>[QS to 200µl]</td>
</tr>
</tbody>
</table>

2. After preparing the hybridization mixture, store the mixture at -20°C.
Hybridization to the Yeast 2.0 GeneChip

After the hybridization mixture has been made, it is denatured at 94C for 5min and injected into the genechip through the injection ports on the back of the chip [see diagram below]. The genechip is then placed in a hybridization oven and the biotinylated target that you prepared is allowed to bind [hybridize] to the sequence specific probes on the array.

Staining the Yeast 2.0 Genechip

The staining procedure involves using a fluidic station designed to accommodate GeneChips. Two solutions are prepared; a strepavidin- phycoerytherin dye (SAPE) and a biotinylated anti-phycoerytherin antibody (BAP). Before placing the GeneChip into the fluidic station, the fragmented cRNA hybridization mixture is manually removed and replaced with buffer.

The chip is placed in the fluidics station and an automatic process of washing and staining occurs for about 2 hours. The procedure starts by first staining with SAPE. The SAPE will bind the biotinylated cRNA that is hybridized to the probes on the chip. It is washed and then stained again with BAP. BAP will bind to the SAPE. The chip is washed again and another SAPE staining is carried out to bind to the BAP. The double staining procedure is employed to amplify the signal intensities. This allows for the detection of low transcript mRNAs that might otherwise not be detected. After staining, the sample is washed and ready for scanning.
Scan the Yeast 2.0 Yeast GeneChip

The GeneChip is scanned after the wash and staining protocols are complete. The chip is placed in the high resolution GS3000 scanner which is equipped with a diode-pumped, frequency-doubled Nd-YAG laser (532 nm) and the 3.5 um beam is scanned back and forth across the glass surface at a pixel resolution of approximately 2.5 um. Fluorescence is detected using a high sensitivity photomultiplier tube [PMT]. This enables high resolution detection of fluorescence from the probe set feature [pixel] that is only 11 um square. There are approximately 1 million probe areas on each chip!
**Images from the GeneChip scanner**

The image on the left is all 1,000,000 probes. The image to the right is zoomed up 20,000 times to show approximately 500 probes. White indicates strong signal [high level of a mRNA in original sample], while dim or no signal indicate little to no mRNA in original sample.

**Bioinformatics**

After collecting and analyzing the image data, the results are archived and burned on to a CD. Because the data files are so large, it is necessary to analyze it using special microarray analysis software. This software can handle tens of thousands of data points at one time and data analysis can be time consuming. Therefore, it is critical that the data meet specific quality control specifications before performing this extensive analysis.
Example Expression Report From MAS 5.0

Date: 04:46PM 01/16/2004
Filename: UVM-test-Yeast.CHP
Probe Array Type: Yeast S98
Algorithm: Statistical
Probe Pair Thr: 8
Controls: Antisense

Alpha1: 0.04
Alpha2: 0.06
Tau: 0.015
Noise (RawQ): 2.970
Scale Factor (SF): 1.367
TGT Value: 150
Norm Factor (NF): 1.00

Background: Avg: 84.82 Std: 2.37 Min: 77.40 Max: 90.40
Noise: Avg: 3.16 Std: 0.15 Min: 2.70 Max: 3.80
Corner+: Avg: 163 Count: 32
Corner-: Avg: 11225 Count: 32
Central-: Avg: 6612 Count: 9

The following data represents probe sets that exceed the probe pair threshold and are not called "No Call".

Total Probe Sets: 14010
Number Present: 5227 37.3%
Number Absent: 8406 60.0%
Number Marginal: 377 2.7%
Average Signal (P): 648.4
Average Signal (A): 32.6
Average Signal (M): 112.0
Average Signal (All): 264.5

Housekeeping Controls:

<table>
<thead>
<tr>
<th>Probe Set</th>
<th>Sig(5')</th>
<th>Det(5')</th>
<th>Sig(M')</th>
<th>Det(M')</th>
<th>Sig(3')</th>
<th>Det(3')</th>
<th>Sig(all)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18S_RRNA</td>
<td>18.3</td>
<td>A</td>
<td>67.8</td>
<td>P</td>
<td>82.9</td>
<td>P</td>
<td>56.33</td>
</tr>
<tr>
<td>ACTIN</td>
<td>2983.0</td>
<td>P</td>
<td>7075.1</td>
<td>P</td>
<td>3976.6</td>
<td>P</td>
<td>4678</td>
</tr>
<tr>
<td>GAPDH</td>
<td>436.2</td>
<td>P</td>
<td>1217.7</td>
<td>P</td>
<td>1692.1</td>
<td>P</td>
<td>1115</td>
</tr>
<tr>
<td>DROS-EIF-4A</td>
<td>4443.5</td>
<td>P</td>
<td>10464.4</td>
<td>P</td>
<td>7337.3</td>
<td>P</td>
<td>7415</td>
</tr>
</tbody>
</table>

Spike Controls:

<table>
<thead>
<tr>
<th>Probe Set</th>
<th>Sig(5')</th>
<th>Det(5')</th>
<th>Sig(M')</th>
<th>Det(M')</th>
<th>Sig(3')</th>
</tr>
</thead>
<tbody>
<tr>
<td>BIOC</td>
<td>331.0</td>
<td>P</td>
<td>155.8</td>
<td>P</td>
<td>243.38</td>
</tr>
<tr>
<td>BIODN</td>
<td>244.2</td>
<td>P</td>
<td>1490.2</td>
<td>P</td>
<td>867.20</td>
</tr>
<tr>
<td>CREX</td>
<td>2554.0</td>
<td>P</td>
<td>5413.1</td>
<td>P</td>
<td>3983.52</td>
</tr>
</tbody>
</table>
Appendix A: Preparing the pre-culture

1] All work must be done in a HEPA hood such as those involved with tissue culture! Always wear gloves that have been rinsed with alcohol!

2] Transfer an entire colony of yeast to a broth culture of 1/2x YPD with or without glucose. It is not important as long as we get some growth as indicated by turbidity at 48 hrs.

3] After incubation, aseptically remove a drop of the culture and examine on the microscope to ensure there is no contaminating bacteria. Remember that bacteria will be 1/5 the size of the yeast. So look for small rods or cocci beyond the yeast.

4] To inoculate the experimental culture to be used in class, aseptically transfer 0.5 ml of this culture to the 1/2x YPD-3xGlu broth 48hrs before day 1. It is advisable to use aerosol resistant pipet tips.

Alternatively:

You may streak a YPD agar plate 4 days prior and then inoculate your experimental culture with a whole colony 48 hours prior to day 1 class.