

Fall 2010: GENE EXPRESSION

ANSC/GENE 626 (edited by. N. Ing 09/08/10)

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Office Hours: Make appointment by email

The objectives of this course are:

- 1. To appreciate and utilize molecular biology in order to assess gene expression**
- 2. To handle biological macromolecules with good techniques to generate high quality data**
- 3. To critically evaluate data and use it to make decisions in experimental plans**
- 4. To communicate experimental purposes, methods, data, and conclusions in written informal and formal formats.**

Class will be held in BICH 243 in Fall semester on Thursdays (lecture from 12:00 to 12:45 p.m.) and Fridays (lab from 1:15 to 3:30 p.m.). First meeting will be in the second week of class. 9/2 and 3. Students should print and read the syllabus and review the protocols independently.

9/9 Lecture 1: Introduction to the course, Safety rules.

9/10 Lab 1. Introduction: Pipetting, Restriction digestion of aromatase plasmid DNA to linearize plasmid for use as a PCR positive control.

9/16 Lecture 2: What is gene expression? What are the functional parts of a gene, an mRNA and a protein? Tripure reagent allows purification of three macromolecules from one tissue sample.

9/17 Lab 2. Extract and precipitate to purify RNA from tissue

9/23 Lecture 3: DNA and RNA structure. Restriction digestion and plasmid analyses.

9/24 Lab 3. Analyze RNA on Northern gels and blot

9/30 Lecture 4: Polymerase chain reaction and uses.

10/1 Lab 4. Clone aromatase cDNA from RNA using reverse transcription and PCR.

10/7 Lecture 5: Molecular cloning for different purposes. Plasmid vectors.

10/8 Lab 5. Analyze PCR products and plasmids on regular agarose gel and purify aromatase cDNA band on low melting point agarose gel

10/14 Lecture 6: Molecular cloning for different purposes II

10/15 Lab 6: Sequence PCR product (in low melt gel) and linearized plasmid (in aqueous solution).

10/21 Lecture 7: DNA sequence: Function and analysis.

10/22: Lab 7: Remove unincorporated nucleotides with spin column, dry and submit for sequencing.

10/28 Lecture 8: mRNA Analyses: a comparison

10/29 Lab 8: Sequence analyses – raw and bioinformatics – of aromatase and “unknown” cDNAs – CLASS MEETS AT WEST CAMPUS LIBRARY Rm ____.

11/4 Lecture 9: Transcription in cells and out

11/5 Lab 9: In vitro transcription for aromatase and “unknown” Northern blot probes and make an acrylamide gel

11/11 Lecture 10: Acrylamide gel electrophoresis for small RNA and protein analyses

11/12 Lab 10: Analysis of the Northern blot probes on acrylamide gels and dot blots

11/18 Lecture 11: Expression of recombinant proteins (Also, perform Northern blot hybridization)

11/19 Lab 11: Northern blot washing and imaging

11/25 and 26 HAPPY THANKSGIVING!!!

12/6 Review of experimental results and laboratory cleanup.

Students can print out the class protocols on my website: <http://animalscience.tamu.edu/ning/lab/courses.htm>. Protocols for the Friday lab should be read by the student before the lecture on Thurs and brought to the lecture and lab classes. Students will be evaluated on the lecture material via pop quizzes, homework problems and, if the instructor deems it necessary, exams (25%). In the lab, grades will be given on preparation and participation (25%), daily lab notebook records of activities (25%) and a written laboratory report in journal paper style (25%) due the Thursday after the last class at 5 PM. A suggested text (An Introduction to Genetic Engineering 3rd ed. by Desmond S.T. Nicholl) has many relevant sections especially for the beginning material. Many reagents and protocols are described in PROMEGA "Protocols and Applications Guide" now available online, <http://www.promega.com/paguide/>.

N.Ing LAB RULES

A. GENERAL

1. Everyone is individually responsible for the experiments. Come prepared by reading protocols and required reading in advance!! Activities will be started immediately, while explanations and discussion sessions will occur as time permits. **COME PREPARED to ASK QUESTIONS**, especially during discussions.
2. Equipment in this and neighboring labs is shared. Know or ask how to use it. Obey user rules, such as signing logs. Leave all equipment in good working order. If there are problems, tell someone so we can fix them!
3. Leave the lab better than you found it. Wash your own glassware, clean up your work area, write the names of reagents that are running out on the "to be ordered" list, etc.
4. **Lab notebooks** are bound volumes, are kept in black ball point pen with numbered, dated pages. They are designated only to that purpose, are labeled on the cover with researcher name, dates of inquiry and laboratory class, and are only removed from the lab with instructor permission. In them, each person describes their activities and observations each day. Each page should be dated. Each data photo or X-ray film should be labeled with initials, date, and identification of gel type, lane contents and dye migration and affixed to a page in the notebook. Record things chronologically. Start a new project on a new page with a description of its purpose. Complete protocols from the class don't have to be written each time in the notebook. Instead, write out the relevant information and refer to the handouts for specifics. For this, you may consider the handouts you receive in lab as references. Use them as such and refer to specific pages (dates) within them. Record your activities and observations clearly, using complete and understandable sentences. Write down the composition of your reagents, including buffers and buffer recipes. A stranger should be able to pick up your notebook and understand why you did an experiment, how you did it, and what results you got. Indeed, they should be able to duplicate your experiments.

KEEP UP WITH NOTEBOOK ENTRIES EVERY DAY...otherwise, data will be lost!

5. All reagents and samples saved must be labeled with black sharpie stating the date; your initials, and **WHAT IT IS**. Professional lab workers use simple sample numbers, such as 1 – 10 that is unique for the date. The full description of the sample is then put in the notebook. Items not labeled sufficiently are worthless and will be discarded.
6. Store things in appropriate places! For plasmids and reactions and buffers, store at – 20C in storage box provided to your group unless otherwise noted. Note storage places in your notebook.

N.Ing LAB RULES (cont'd)

B. SAFETY IS THE #1 PRIORITY

1. The only safety activity not strictly enforced is wearing safety glasses: this is a good idea but is not mandatory. Wearing a lab coat is mandatory and wearing gloves will become a habit (see below).
2. Working with open flames and hazardous chemicals have strict safety protocols - ask for them and follow them.
3. Working with radioactivity is a privilege, not a right. Workers must monitor for contamination, before, during, and after the procedure. Radiation safety training is required. WE RUN A CLEAN LAB.
4. Because we work with HAZARDOUS SUBSTANCES, there is NO EATING, DRINKING, SMOKING, or APPLYING MAKE-UP in the lab.
5. Garbage must be disposed of properly. Glass and sharps, biohazard, chemical, and radioactive waste must be separated from the rest.

C. GOOD LAB TECHNIQUES

1. ICE IS NICE! Get it in BioBio Room 229. Work on it unless otherwise directed. It slows degradation of macromolecules.
 2. Many reagents settle on storage, so mix them! All frozen solutions need to be thawed and mixed before using.
 3. ENZYMES DO OUR WORK. They are stable as glycerol solutions at -20°C. Keep them in the freezer as much as possible. Only remove them in -20°C blocks. DO NOT WARM ENZYME STOCKS! When pipetting small amounts of viscous solutions like enzymes, check loaded pipet tip and evacuated one to assure that enzyme got into the reaction. After addition, mix reaction solution gently but thoroughly: can pipet total volume up and down OR vortex gently and flash spin in microfuge to return reaction to the tube bottom.
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ADA Statement, Copyrights, and Plagiarism

The Americans with Disabilities Act (ADA) is a federal antidiscrimination statute that provides comprehensive civil rights protection for persons with disabilities. Among other things this legislation requires that all students with disabilities be guaranteed a learning environment that provides for reasonable accommodation of their disabilities. If you believe you have a disability requiring an accommodation, please contact the Dept. of Student Life, Services for Students with Disabilities in Room 126 of the Koldus Bldg. or call 845-1637.

Copyrights

The handouts used in this course are copyrighted. By handouts", I mean all materials generated for this class, which include but are not limited to syllabi, quizzes, exams, lab problems, in-class materials, review sheets, and additional problem sets. Because these materials are copyrighted, you do not have the right to copy the handouts unless I expressly grant permission.

Plagiarism

As commonly defined, plagiarism consists of passing off as one's own ideas, words, writings, etc., which belong to another. In accordance with this definition, you are committing plagiarism if you copy the work of another person and turn it in as you own, even if you should have the permission of that person. Plagiarism is one of the worst academic sins, for the plagiarist destroys the trust among colleagues, without which research cannot safely be communicated.

If you have any questions regarding plagiarism, please consult the latest issue of the Texas A&M University Student Rules, under the section "Scholastic Dishonesty."

BIO 243 Lab Safety Precautions

1. ABSOLUTELY no food or drink in the lab. This includes gum.
2. Always wear gloves when working with hazardous compounds: ethidium bromide, radioactive compounds, acrylamide and organic compounds. Wear gloves to protect reactions from contamination from you (RNase, DNA) when working with RNA and PCR.
3. Always wear UV safety glasses when using UV illumination.
4. Use special care when working with open flames. Don't forget to turn off gas after use.
5. Clean up spills immediately. Notify instructor if hazardous compounds are spilled.
6. Discard organic solutions in appropriate waste bottles in fume hoods.
7. Discard Ethidium bromide waste in correct container.
8. All culture medium and labware used for bacteria needs to be autoclaved or add Chlorox to liquids to final concentration of 5 or 10 % before disposal and mix and let sit 5 min.
9. All sharps (including broken glass, needles and razor blades) should be disposed in clearly marked containers, not in the general trash.
10. If you have questions about anything, ASK!
11. Lab coats are required!

I have read and will follow the above safety rules.

Signed _____
Date _____

LAB REPORTS -

by Linda Guarino and Nancy Ing

TITLE - This is the most important part of a manuscript. A reader begins here, and will also finish here if the title does not promise a subject of interest to him/her. A good overall rule is to use the fewest possible words that adequately describe the contents of the paper. But, do not sacrifice words for specific information. For example 'DNA cloning' is a short title, but it is too general. A popular trend in recent years is to publish papers where the title is a complete sentence that summarizes the major conclusion of the manuscript. Personally, I prefer titles that describe the work, not the results.

ABSTRACT - An abstract is a mini version of the paper. It should provide a brief (less than 250 words) summary of the major points of the manuscript. The abstract should state the objectives, describe the methodology used, summarize the results, and state the principle conclusions. The abstract should be written in the past tense, because it refers to work done.

INTRODUCTION - First of all, state the nature and scope of the problem investigated. Review the pertinent literature (**NOT NECESSARY FOR THIS CLASS**). Describe the method of the investigation. State the principle results. State the principle conclusions suggested by the results. The first two parts should be in present tense, while comments relating to the present study should be in past tense.

METHODS - The methods section should expand upon the description of the methodology that was presented in the abstract. The order of presentation is usually chronological (methods used in initial stages of the study are presented first). However, sometimes it makes more sense to group similar methods into sections, even though they were not used at the same time. Due to space limitations in journals, methods are not usually described in detail if they have previously been published. If a scientist uses a protocol that is identical to one previously described, he/she would state 'The DNA was prepared according the procedure previously described (reference). If there were minor differences, he/she would state 'according to the procedure of (ref.) with minor modifications' and then describe the modifications. In this class, you may assume that the class protocol has been published. Therefore you don't need to give the details, but you need to describe the general strategy. For example, you should say 'the DNA was purified by the alkaline lysis procedure as previously described' not 'the DNA was purified as previously described'. In addition to the class protocol, you could also reference the Cloning manual or the Promega manual. The methods section should be written in past tense.

RESULTS - The results section is a presentation of the data. It should not repeat the methods given in the previous section. Each figure should be referred to here. The results section should be written in past tense.

DISCUSSION - The discussion should put the results into perspective. Discuss the results without recapitulating the results section. Show how your results and interpretations agree. State your conclusions clearly, and summarize the evidence for each conclusion. Selection of correct tense is more difficult in the discussion than in the other sections. Your own work should be described in past tense. If reference is made to published work, it should be in present tense.

REFERENCES - Only need to cite the class protocols and any other sources of material...no need for literature review so these are very few.

FIGURES and FIGURE LEGENDS - Present the important data in figure form, raw data if possible for this class. Figures should have complete legends - so that they can be understood without reading the rest of the paper. Figure legends begin with a title for the figure. Then the legend should completely describe what the figure shows. Therefore, the figure legends may contain information that is redundant with information in the Methods and Results sections. Figures and figure legends may be nested in the paper or placed at the back.

Many students have asked about length. The best rule that I can give you is that it should be long enough to convince me that you have learned something. However, I have a short attention span, and if your paper is very long and verbose, I may lose interest before I decide whether or not you have learned anything.

Suggested reading:

Day, R. A. 1988 How to write and publish a scientific paper, 3rd ed. Oryx Press, Phoenix.

9/10/10

GENE EXPRESSION LAB 1

A. Practice pipeting: The better you are at pipeting, the better your results will be!

{Can do 1, then 2 then 3 or 1, then 3 then 2.}

1. Read instructions for Use of Micropipettors (below).
2. Determine densities of liquids:
 - Tare a 1.5 ml test tube on the balance.
 - Weigh 1000 ul of water two times.
 - Repeat with Isopropanol.
 - Determine the densities of these liquids and write them in your notebook.
3. Measure the volumes of the unknown samples provided in the 1.5ml test tubes labeled "A", "B", and "C". Record in your notebook.
4. Check your results with an instructor.

Use of Micropipettors

1. Choose the correct pipet. For volumes:
 - 1-20 μ l P20
 - 20-200 P200
 - 200 - 1000 P1000
2. Set the desired volume by holding the pipetman in one hand and turning the volume adjustment knob until the correct volume shows on the indicator. For best precision, always approach the desired volume by dialing downward (at least one-third revolution) from a larger volume setting.
3. Attach a new tip to the shaft of the pipet. Press tip on firmly to ensure airtight seal. Choose the correct tip.
 - P20 yellow tip
 - P200 yellow tip
 - P1000 blue tip
4. Depress plunger to first positive stop. Hold pipetman vertically and immerse disposable tip into sample liquid 2mm.
5. Allow the push button to return slowly to the up position. Never permit it to snap up.
6. Wait 1 or 2 seconds to ensure that the full volume of the sample is drawn into the tip.
7. Withdraw tip from the sample liquid. Wipe the sides of the tip on the sides of tube to remove any remaining liquid.

8. To dispense the sample, place the tip end against the side wall of the receiving vessel and depress the plunger slowly to the first stop. Then depress the plunger to the second stop to expel any residual liquid in the tip.

9. With the plunger fully depressed, withdraw pipetman from the vessel. Then allow the plunger to return to the top position.

10. Discard tip by depressing the tip ejector button. A fresh tip should be used for each sample.

B. Restriction digestion of aromatase plasmid DNA.

NOTE: Restriction Enzyme digestions usually contain 1-2 ug DNA and 5-12 units of enzyme in 20 ul. Working buffer strength is always 1X. Use the buffer supplied with the enzyme (optimized) if it is a single enzyme cut. If using two enzymes, use a buffer compatible to both that gives at least 75% effectiveness! There is a table in the Promega book that will help you choose.

1.. Plan a 20 ul restriction digest for the unknown plasmid containing a cDNA that you were given. Plasmid concentration is also given. From that calculate the volume of plasmid you will add to the reaction to get 0.5 ug:

_____ ul H₂O (to make reaction 20 ul final volume)

2 ul (which??????) 10X Buffer

_____ ul (0.5 ug) aromatase plasmid DNA

1 ul Xba I restriction enzyme

20 ul TOTAL reaction volume

NOTES ON USING ENZYMES: These are the expensive workhorses of molecular biology. Your success depends on their function. Therefore, they should be handled with care: KEEP THEM AT -20°C at all times by setting up the entire reaction prior to retrieving them from the freezer. Keep them in the -20°C blocks while pipeting. They are in glycerol, so are viscous. Visually check that enzyme is picked up in the pipet tip and delivered into each tube. Ensure their purity by using new pipette tips each time!!!! After addition, mix the enzyme into the reaction by gently pipeting the whole reaction up and down, or by vortexing gently and flash microfuging.

2. Check calculations for the reaction with the instructor, then set up the digestion.

3. Incubate at 37oC for 1 h.

4. Store reactions at -20oC.

DISCUSS as a class: DNA sequence and structure using human nucleotide models

1. Make a six base DNA single strand (random sequence)...Identify 5' and 3' ends. What type of bonds exist between bases?
2. What is the chance that a specific six base sequence will occur?
3. Make a complementary DNA strand hybridized to the one in #1. How are the strands oriented to each other? What type of chemical bonds exist between the DNA strands?
4. Reverse the 5' to 3' direction of the complementary strand. What bonds (if any) did you have to break to do this? How is this strand related to the initial strand?
5. Make an EcoR I restriction enzyme site. What bonds does the enzyme cut? What makes the ends "sticky"? Do the ends have 5' or 3' overhangs or one of each?

Now restore the restriction enzyme site integrity with T4 DNA ligase.

RNA ISOLATION FROM MAMMALIAN TISSUE

TODAY you will extract RNA from tissues for analysis on a Northern blot and generating cDNAs for cloning. The amount of RNA obtained is measured by sample absorbance at 260nm. To assess RNA quality, the RNA preparation is analyzed on a denaturing gel. This is transferred to a membrane ("Northern blotting") for hybridization with a probe to sensitively detect a specific mRNA.

A. Preparing Materials for RNA Work

Avoiding degradation of RNA by RNase

You need to read about RNase (below and http://www.ambion.com/techlib/tb/tb_159.html), a ubiquitous enzyme that efficiently destroys RNA. Primarily, this will serve to make you paranoid and do neurotic things, like wear gloves all the time. Although working with RNA is similar to working with DNA, many RNA experiments fail miserably because of RNase, so know this enemy!

In biochemistry, RNase is the model of an enzyme that will not die: not in an autoclave or even after dehydration (by alcohol, etc.). As soon as it returns to a water environment between room temperature (R.T.) and 37°C, it chews again. It is an enzyme of all living things and is important in keeping RNA turnover high so cells aren't overwhelmed by too much RNA and so new expression of genes tightly regulates cell function.

The best way to beat RNase is to avoid it. Work with the cleanest reagents and lab-ware. Things that aren't handled by or contaminated by living things are generally RNase-free; e.g. paper towels. Test tubes and pipettes don't have to be sterile but should be used from freshly opened packages. Then protect packages from dust and fingers by resealing packages and storing in cabinets. Glassware is reserved similarly: wrapped and stored away from general use. Equipment like Pipetmen and Gel apparatus for RNA are reserved for this use and are NOT USED WITH RNase!

Solutions are made with water of the highest purity. Dry chemicals are shaken out of containers: residual amounts are discarded. Nothing dirty (like spatulae or pipettes) is introduced into chemical stocks: solutions or powders.

All solutions are treated with 0.1% diethyl pyrocarbonate (DEPC). This oily liquid is added. The solution is shaken vigorously until foamy (aerobic workout). The solution is incubated 37°C or room temperature overnight to allow the DEPC to covalently attack RNase. The solution is then autoclaved to destroy DEPC (which also attacks RNA) and to sterilize to prevent growth of undesirables. Exceptions to this solution preparation protocol are 20% SDS (nothing grows in this) and Tris solutions (which DEPC attacks, too). NOTE: DEPC treatment can only correct a low level of RNase contamination!

You must start clean!!!

Make 1 li DEPC-H₂O per person

1. In a clean (RNase-ZAPped) 1 liter bottle add nanopure H₂O to 1 liter level for DEPC-H₂O.

2. Add 1ml DEPC per liter.

3. Shake till foamy for 10 sec.

4. Put in 37°C incubator O/N to allow DEPC to work optimally.

These will be autoclaved for 40 min to destroy DEPC before use.

[Autoclave the DEPC-H₂O to destroy DEPC and prevent any growth in solutions that might introduce RNase. NOTE on Autoclaving: Need 35 to 40 min. sterilization time for 1 liter. 20 min. for 500ml. Use "liquid" cycle and keep caps loose]

B. RNA Extraction

Read pertinent parts of

http://www.roche-applied-science.com/proddata/gpip/3_6_6_2_19_1.html

NOTE 1: Tripure has phenol and guanidine salts in it...both are caustic and burn skin!!!
Be careful! Wear safety glasses!!!

NOTE 2: CHCl₃ (Chloroform) dissolves things like styrofoam and polystyrene - use glass graduated pipets and polypropylene 15 & 50 ml tubes.

You know how to fight RNase to keep materials clean. GUESS WHAT! RNase is in all living systems including the one from which you'll purify RNA. So all RNA preparers begin with the realization that their worst enzyme enemy is present in the sample. In the cell, RNA is compartmentalized away from RNase so many tissues are OK for harvesting for RNA if kept coolish even up to 24 h after collection (of course, faster may be better). But freezing breaks intra-cellular membranes, mixing RNA with RNase. Therefore, fresh tissues are kept cool while mincing and weighing, then are put in a 1.5 ml polypropylene tube snap frozen in liquid N₂. They are stored at -80°C. They may store well for 6 months but usually not for more than 1-2 years. This is dependent on them avoiding thawing, too. So the TWO MAIN POINTS about tissue collection are to SNAP FREEZE and KEEP at -80°C until use within 1 year. NOTE: You can't snap freeze things much bigger than 25 mm³. I mince to about 8 mm³. Many people use RNALater to store tissues for later RNA preparation and have good success.

Tissues vary with RNase content and amount of connective tissue present, so RNA yields vary in quality and amount. RNA extraction from cultured cells results in very high quality RNA, usually.

RNA Extraction from tissue with Roche TriPure reagent (contains phenol! see NOTE 1 below)

Each student will do 2 RNA preps (one from pig endometrium, one from horse testis). Label all tubes needed NOW!

1. Homogenize 0.25 – 0.5 mg tissue (frozen or fresh) in 5 ml room temperature ("RT") Tripure solution in a 15 ml polypropylene tube. Use three 15 sec bursts at 70% power. Rinse probe in tripure (do a mock homogenization with Tripure and no tissue) between dissimilar samples.
2. Incubate RT 5min. During this time, transfer the contents equally into 4 - 1.5 ml tubes.
3. Add 250 ul chloroform to each 1.5 ml tube using a P-1000. Mix by vortexing or shaking vigorously 15 sec.
4. Incubate RT 5 min.
5. Centrifuge 15 min at 10,000 rpm at RT or 4oC in a microfuge.
6. Transfer upper phase to four clean 1.5 ml tubes with transfer pipet. AVOID THE INTERPHASE!!!! Discard lower phase and interface in phenol waste container.
7. Precipitate RNA by adding an equal volume of isopropanol. Mix by inverting tube. Incubate RT 5 min
8. Centrifuge at 10,000 rpm for 10 min at RT.
9. Wash pellet in 75% EtOH (make 10 ml with 100% EtOH and DEPC H₂O). This means to discard the supernatant, add the supernatant volume of wash (75% EtOH), vortex, microfuge 5 min, and discard supernatant. The purpose is to wash salts out of the RNA pellet, which should not dissolve during the procedure.
10. Air dry pellet briefly after spin. You can wipe the sides of the tubes with Kimwipes, but stay away from the pellets! Do not dry totally or you will not be able to solubilize RNA easily!!!
11. Dissolve the four similar pellets each in 25 ul 1 mM Na citrate Buffer/pH 6.4 or TE buffer (10 mM Tris, 1 mM EDTA pH8).
(Heat in 70oC block and vortex hard and repeatedly over 15 minutes.) Pool so that you have a 100 ul sample for each RNA prep.
12. Use Nanodrop in Kleberg 414 to measure RNA concentration (A₂₆₀) and estimate purity (proteins and phenol absorb at A₂₈₀) according to instructions.
13. If RNA concentration is 4 ug/ul or more, put 20 ug samples of each RNA preparation into a new labeled tube (label "C" for endometrium and "D" for testes). If RNA concentration is less than 4 mg/ml, just put 5 ul in the tube and note how many ug of RNA is in that sample in your notebook.
14. Store RNA at -80oC

{**RNA STORAGE:** Store at 4oC during sample use (this class). For storage over 1 week, can store at -80oC. For longer storage, add 3 volumes of 100% ethanol and store at -80oC.}

GENE EXPRESSION LAB 3 ning 9/24/10

RNA ANALYSIS ON NORTHERN GELS/BLOTS

[See NorthernMax (Ambion) protocol]

TODAY we will electrophorese RNA from tissue and controls on a Northern gel that will be blotted to a nylon membrane for probing with an antisense complementary RNA (cRNA) probe.

Be clean – clean and protect apparati from Rnase contamination- work on clean diapers!
Use all DEPC reagents, gloved hands and aerosol barrier tips.

I. Your gels have already been poured this morning for you as described below: **2 people per gel.**

1. Rinse beaker/flask well with house-distilled water. Melt agarose (0.8 g) in 72 ml DEPC-H₂O in an RNASE-Free glass bottle or beaker. (Bring to a boil in microwave oven and mix by swirling: repeat 2 to 3 times).
2. Cool to 70oC.
3. In a fume hood, add 8 ml 10X Denaturing Gel buffer (formaldehyde and MOPS/pH 7.0, NaAc, and EDTA) and pour into RNase-free gel mold with the ends taped. Use two thin combs.

II. Control RNA samples prepared for you – ready to heat and load

A. Positive control RNA in Northern Sample Loading buffer in 25 ul.

B. An “unknown” plasmid DNA template for in vitro transcription of a cRNA probe. This will serve as a positive control for electrophoresis, transfer, hybridization, and development of signals. A 10 ng sample is in 25 ul of Northern Sample Loading buffer (+ EtBr)

III. You need to prepare your RNA samples C (endometrium) and D (testis):

- A. If sample is less than 5 ul, add DEPC-treated water to make volume 5 ul.
- B. Add 15 ul of Northern Sample Loading buffer (+ EtBr) RNA tubes for gel..

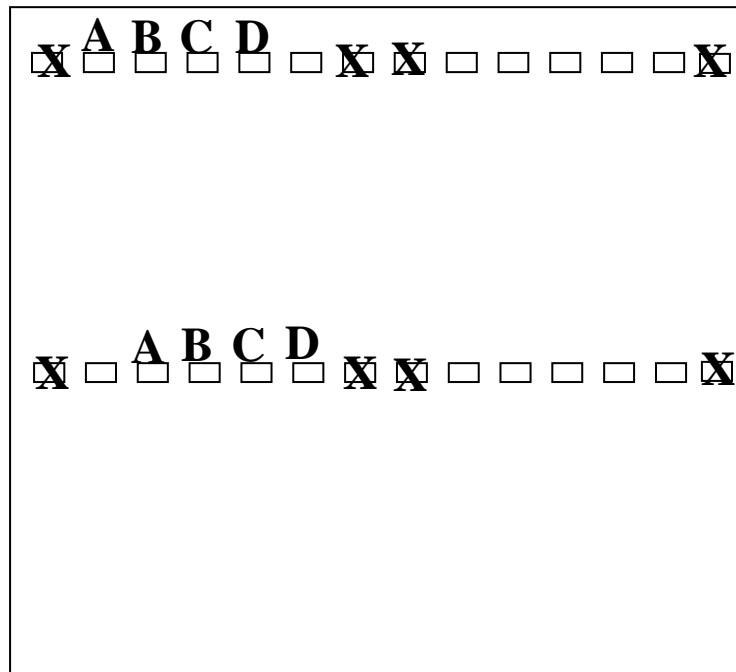
III. Load and Run the Northern gel.

A. All RNA samples should be heated at 68oC for 10 min. and then chilled on ice prior to loading.

B. Clean a gel rig with RNaseZAP! And fill it with 1 li of 1X Gel Running Buffer. Remove tape from gel and submerge it before removing the combs.

C.. Load samples A to D onto the gel under 1X Gel Running Buffer (dilute the 10X stock from the kit with your DEPC-H2O bottle, cover the gel with buffer to about 0.5 cm depth). We won't use RNA markers...we can use the internal ribosomal RNAs (rRNAs) as internal markers in samples A and C and D.

One student load 10 ul volumes A-D on top and bottom of left side of gel, then the other student load similarly on left side. Do not use far outside lanes or the two most central lanes. Left and right halves of gels should contain two replicate sets of samples. Prior to hybridization, top and bottom halves will be separated and probed with different probes. Diagram below is after first student loaded samples.



D. Run at 100 volts until dye front reaches bottom or sufficient separation occurs (1 to 1.5 h); can peek at gel progress with hand-held short wave UV lamp. Prepare materials for blotting during this time!!!!!!!!!!!!!!

E. Take photo on UV box alongside a fluorescent ruler.

IV. Northern transfer to Brightstar nylon membrane

(Note: This is different than that in the NorthernMax kit instructions)

1. Cut wicks (2 pieces 14 cm X > 30 cm), blotting papers (3 pieces 11 X 14 cm), and nylon membrane (1 piece of 11 X 14 cm) wet in transfer buffer as directed and pictured in the diagram. Also need paper towels cut in half to be about 14 X 11 cm....need a 4 inch tall stack of these.

2. Assemble an upward capillary transfer as instructed in the diagram provided. Align the the blot with the gel especially at the tops. You can use the rig you ran the gel in to do the transfer to nylon. Allow the transfer to continue with 500 ml of 10X SSC until the next lab.

QUESTION to be answered in your notebook: What are the compositions of your northern gel, running buffer and sample buffer used today?

GENE EXPRESSION Lab 4 ning 10/1/10

Northern Blot crosslinking and RT-PCR amplification of a cDNA

Note: Can program the PCR machine anytime during the lab.

A. REVERSE TRANSCRIPTION:

Today each student will reverse transcribe their two RNA preps with random hexamer primers to make cDNA, which is capable of being amplified by PCR. Each student should set up four reverse transcription reactions: two will be the real reactions, one for each RNA prep containing reverse transcriptase enzyme (#1 and #2). The other two will be mock reactions, identical to #1 and 2 but DEPC-treated water used instead of enzyme (#3 and #4).

Label four sterile 500 ul tubes. From the Superscript II Reverse Transcriptase protocol:

1. For each 20 ul RT reaction, combine in a tube:

Random primer	0.25 ug/ul	1 ul
1ng – 5ug of total RNA		X? ul
dNTP mix (10mM each)		1 ul
Sterile, distilled water		to 12 ul total

2. Heat tubes to 65 degrees C for 5 minutes, quick chill on ice, and flash spin, and add to each;

4 ul 5X First Strand buffer (with kit)

2 ul 0.1M DTT (with kit)

1 ul Rnasin (Promega)

3. Mix contents of the tubes gently. Incubate at 25 degrees C (room temp.) for 2 minutes.

4. Add 1 ul of Superscript II enzyme to reactions #1 and 2 and mix by pipetting up and down. To reactions #3 and 4, add 1 ul H₂O and mix similarly.

5. Incubate tubes at 25oC (room temp.) for 10 minutes. Then incubate reactions at 42oC for 50 minutes.

6. Inactivate the enzyme (all tubes) by heating at 70 degrees C for 15 minutes. Store in -20°C.

B. Northern Blot crosslinking

1. Remove all papers from the Northern transfer but keep the gel and blot/filter together.

2. IMPORTANT: Mark well positions on the blot with a sharp pencil or a black Sharpie marker. Also write initials and date. Put these marks on filter's back.

NOTE: Keep the blot RNA-side-up during subsequent handling and keep it in a *clean, labeled* container.

3. In a Tupperware container with lid, rinse agarose particles from filter in 2X SSC with vigorous agitation for 30 sec.

4. Put blot on plastic wrap and view it on a UV box. On the side of the blot, mark positions of 28S and 18S rRNAs if those bands were visible in your tissue RNA lanes.
5. Look at the gel on the UV box. Did all of the RNA transfer out?
6. Place wet blot on top of Whatman paper saturated with 2X SSC - all on top of plastic wrap.
7. UV crosslink RNA to nylon (use Stratalinker in energy mode: 120,000 ujoules).
8. Store the blot between clean paper towels in a labeled Ziploc bag in the -20oC freezer.
9. Clean, dry and put away your labware used for blot transfer.

C. PCR AMPLIFICATION OF A cDNA

Today each student will amplify an aromatase cDNA fragment from a specific mRNA (unknown, will be identified by sequence analysis later) among the reverse transcribed cDNAs in the reverse transcription reaction from last week. One negative control is water (no DNA template). This control is a must for all PCR runs. More rare negative controls are the mock reverse transcription reactions that lacked reverse transcriptase enzyme. A positive control is the linearized plasmid containing the cDNA from Lab 1.

For PCR you should be clean! This means do not contaminate the reactions with your DNA (on your hands and on pipettors). Wear gloves and use aerosol barrier tips to optimize cleanliness.

Label 7 thin-wall 200ul PCR tubes around the necks of the tubes. Primers will be provided at concentrations of 70 ng/ul

Each student will set up 6 - 25 ul reactions, so make a Master Mix of common reagents for 7 reactions (due to the Law of Disappearing Volume) in a 500 ul tube:

	<u>In one reaction:</u>	<u>In Master Mix for 7 reactions:</u>
H2O	18.68 ul	131 ul
10X Taq Buffer	2.5 ul	17.5 ul
2.5 mM dNTPs	2 ul	14 ul
Primer 1	0.5 ul	3.5 ul
Primer 2	0.5 ul	3.5 ul
Taq enzyme	0.125 ul	0.875 ul
TOTAL	24 ul	

Pipet 24 ul of Master Mix into each labeled PCR tube.

Then add the unique DNA template (1 ul) that will be amplified from during the PCR:

1. Water (no DNA template)
2. Reverse transcription reaction #1
3. Reverse transcription reaction #2
4. Mock reverse transcription reaction #3

5. Mock reverse transcription reaction #4
6. Linearized plasmid DNA containing the aromatase cDNA – Made in the first lab.

Place the tubes in the PCR machine and run a program for 35 cycles of:

94°C denaturation for 30 sec,
45°C annealing for 1 min, and
72°C extension for 1 min.

The machines will run the PCR then the tubes will be stored at 4°C until the next lab.

Questions:

1. What are the DNA sequences of the primers used today?
2. What are the compositions of the buffers used in enzymatic reactions today?

GENE EXPRESSION Lab 5 ning 10/8/10

Analyze PCR products on agarose gels and cut a cDNA band for an in vitro transcription template.

MAKE a 1% agarose gel for DNA analysis and a low melting point agarose gel for PCR band purification: See appendix. Need 1 gel per 2 to 4 students.

PREPARING SAMPLES AND RUNNING THE GEL

1. In clean tubes, one for each PCR reaction, combine 25 ul of PCR product with 2.5 ul of 10X DNA Loading buffer containing two tracking dyes (bromphenol blue and xylene cyanol). Mix well.
2. Submerge the gel under 900 mls of 1X TAE buffer in the running chamber.
3. Load the 12 ul PCR samples plus Loading buffer alongside a sample of 10ul (1 ug) Lambda HindIII EcoR I DNA markers on each of the two agarose gels. Record your loading order (including lane location on gel) in your notebook!
4. Run the gels at 100 V for 45 min.

FOR THE ANALYTICAL AGAROSE GEL:

1. After the gel run, visualize bands with short wave UV light and take a picture with the camera imaging system.
2. By comparing the bands in PCR sample lanes with Lambda DNA markers or standards, one can describe
 - a. The molecular size of the PCR products and
 - b. The amount of DNA in PCR product.

For (a), compare migration distance to that of Lambda standard fragments - estimate bp size.

For (b), compare brightness of bands to those of Lambda fragments - estimate ng of DNA in the band....use the provided description of the marker bands.. Divide by ul of original DNA sample to get its [DNA]. This is often more reliable DNA quantitation by A260 measures!!!

For products of at least one lane, describe these aspects of DNA fragment analysis. Write a text description of PCR/gel results.

NOTE: On the periphery of the printed picture (not on the data itself) MARK DYE POSITIONS ON GEL PHOTOGRAPHS. Label all photos completely! Name, date, identity of gel and lane components!!! This should be taped into your notebook.

FOR THE PREPARATIVE LOW MELTING POINT AGAROSE GEL:

1. From the photograph of the analytical gel, choose a PCR band from one of your experimental reverse transcription reactions to cut out for sequencing and to use as a template for in vitro transcription.
2. Label a clean tube and get a clean gel cutting instrument or pipette tip.
3. In the prep room with lights off, visualize the band with hand held long wave UV light source while everyone nearby is wearing UV protective goggles.
4. Cut the DNA band with little or no surrounding gel and put in the tube for storage at -20oC. After you have cut the band can visualize and photograph the low melt gel as you did the analytical gel (above.)

Sequence the purified PCR band (in low melt gel) and an unknown plasmid (in aqueous phase)

The procedure is called Cycle Sequencing. It is basically dideoxynucleotide sequencing or Sanger sequencing with Polymerase Chain Reaction using only one primer. The primer is “T7” which is the T7 promoter sequence: TAATACGACTCACTATAGGG.

Each student will set up and run two sequencing reactions, one for the aromatase cDNA band purified in gel and one for an unknown plasmid (to be identified by sequencing).

1. Program the PCR machine for

30 cycles of: 96°C 20 sec

50°C 10 sec

60°C 4 min

2. In a labelled 200 ul thin-walled PCR tube, combine:

2 ul Big-Dye mix* (Applied Biosystems product)

___ul DNA template (400 ng)

___ul “T7” PCR primer (10 pmol)

___ul ddH₂O (to final volume 7 ul)

3. Mix well and place in a thermal cycler

4. Run the PCR

5. Either set thermal cycler in a cold room so samples will be held at 4°C or retrieve tubes and store at 4 °C later today or tomorrow.

*Q.: What’s in the mix?

Can Watch VIDEO: Dideoxy Sequencing reaction

GENE EXPRESSION Lab 7 ning 10/22/10

Removal of Unincorporated Nucleotides from PCR sequencing Reactions and submitting for automated sequencing:

A. To separate the labeled DNA from unincorporated labeled nucleotides, a spin column is used. These can be made, as described below, or purchased. Centrifugation forces the fluid through the gel matrix of the column. The small nucleotides get hung up in the beads, while the large DNAs fall through and are collected in a tube below the column.

1. Get a TAMU Biology Gene Technologies Micro Bio-Spin Chromatography column in water (not TE) and collection tube (2 ml).
2. Invert the column sharply several times to resuspend settled gel and remove bubbles. Snap off the tip and place column in 2 ml tube. Now remove top cap. Allow buffer to drain by gravity and discard it.
3. Centrifuge for 1 minute in a tabletop IEC centrifuge at **1000 rpm** and discard buffer.
4. Increase the PCR volume to 30 ul with water. Load the PCR sequencing reaction in a 30 ul volume on the center of the gel column with a clean tube (1.5 ml) as collection tube.
5. Centrifuge for 1 min at 1000 rpm. You may have to cut the tube cap off so label the tube body.
6. Open the tube cap. These samples will be speed-vacuum dried, then submitted to Biology Gene Technologies core lab for running on an automated DNA sequencer.

Can Watch VIDEO: DNA Sequencing Electrophoresis

GENE 626 - Lab 8 10/29/10

DNA SEQUENCE ANALYSIS – CLASS MEETS AT WEST CAMPUS LIBRARY ROOM

Sequence analysis and similarity searches in GenBank at NCBI*

*** Q: What does NCBI stand for?**

DNA sequencing provides the ultimate description/information about the DNA. From DNA sequence you will learn what fragment of aromatase mRNA you cloned and WHAT GENE PRODUCT IS IN THE UNKNOWN PLASMID.

Since DNA sequence is information dense, one really only needs 50 - 100 bases of good data to identify the cDNAs...choose your best sequence from the sequencing chromatogram. The sequence reads from left to right as 5' to 3' and will begin about 50 bases 3' from the primer binding site. The aromatase cDNA is not cloned into a plasmid vector so the sequence will begin within the cDNA. The unknown is in the plasmid pTRI from AMBION so you may get some plasmid vector sequence.

1. Make an electronic file of 50 to 100 bases of good cDNA sequence from each reaction. Make sure you enter the sequence in the 5' to 3' direction. (If you do not, the reverse sequence actually bears no relation to the correct one except for having the same overall nucleotide composition.)

2. Copy one sequence and search for similar sequences in GenBank with the BLAST software at the Genbank website (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>).

3. Choose "BLAST Sequence similarity searching." Choose "nucleotide BLAST."

4. Paste your sequence in the box. Choose Database "Other nr" the nonredundant nucleotide database. "Optimize for" choose "Somewhat similar sequences" since we are using non-human and non-mouse sequences. Click on the BLAST button on the bottom

{ You can enter the sequences of the cDNA in "FASTA Format": the first line can contain a name of the sequence and other information preceded by a ">" . Hit "enter" and start entering sequence only on the next line. }

If you are working during the weekday, sometimes the search takes a while. You can e-mail the results to yourself to save yourself from a long wait. Or you can wait and get the results from the website.

Sample BLAST results are:

BLASTN 2.0.3 [Nov-14-1997]

Reference: Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schälffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", *Nucleic Acids Res.* 25:3389-3402.

Query=
(63 letters)

Database: Non-redundant GenBank+EMBL+DDBJ+PDB sequences
323,678 sequences; 623,222,891 total letters

Searching done

Sequences producing significant alignments: (bits) Score E Value

```
emb|X06624|OCPRR3 Human progesterone receptor mRNA 3'-flank 109 3e-23
emb|X51730|HSPREC Human mRNA and promoter DNA for progesterone r... 48 9e-05
gb|U91328|HSU91328 Human hereditary haemochromatosis region, his... 36 0.36
gb|U09532|HFU09532 Heterodontus francisci clone HFB15 T cell rec... 34 1.4
dbj|AB004541|AB004541 Dugesia japonica mRNA for serotonin recept... 34 1.4
emb|X59371|SCURAL Yeast URAL gene for dihydroorotic acid dehydro... 34 1.4
emb|X75951|SC6ORF S.cerevisiae URAL, SAC1, RSD1 and TRP3 genes a... 34 1.4
emb|Z28215|SCYKL215C S.cerevisiae chromosome XI reading frame OR... 34 1.4
```

```
>emb|X06624|OCPRR3 Human progesterone receptor mRNA 3'-flank
Length = 3566
Score = 109 bits (55), Expect = 3e-23
Identities = 62/63 (98%), Positives = 62/63 (98%), Gaps = 1/63 (1%)
```

```
Query:      1 attattgaagtaagctatgtcttaccataactatttcataccatttaagtgaggatttt 60
            |||
Sbjct:     273 attattgaagtaagctatgtcttaccataactatttcata-ccatttaagtgaggatttt 331
```

```
>emb|X51730|HSPREC Human mRNA and promoter DNA for progesterone receptor
Length = 5003
Score = 48.1 bits (24), Expect = 9e-05
Identities = 49/56 (87%), Positives = 49/56 (87%), Gaps = 1/56 (1%)
```

```
Query:      8 aagtaagctatgtcttaccataactatttcataccatttaagtgaggatttttaa 63
            |||
Sbjct:     4512 aagtaaactatatcttatccatattatttcata-ccatgtagggtgaggatttttaa 4566
```

```
>gb|U91328|HSU91328 Human hereditary haemochromatosis region, histone 2A-like protein gene,
hereditary haemochromatosis (HLA-H) gene, RoRet gene, and
sodium phosphate transporter (NPT3) gene, complete cds
Length = 246282
Score = 36.2 bits (18), Expect = 0.36
Identities = 21/22 (95%), Positives = 21/22 (95%)
```

```
Query:      1 attattgaagtaagctatgtct 22
            |||
Sbjct:     217521 attattgaagtaagcaatgtct 217500
```

JUDGING THE BLAST RESULTS:

5. To determine if the sequences listed are really related, look at the E score. It should be less than 0.05 to have a 95% confidence interval that the sequences (query and subject) are truly related. The top hit in the example has very high confidence because E value is very small (9×10^{-5}).
6. The best "hits" should also be logical: You sequenced a cDNA clone so we are looking for a hit on a known mRNA or an exon of a gene. Therefore, note the sequence position numbers of the Subject sequence that your entered sequence (Query) aligns to. On the top hit, click the GenBank report and look at Features, make sure the region of the Subject sequence relates to an mRNA sequence. Also, look at where the cDNA aligns to (sequence position numbers of the subject sequence) in the GenBank report.
7. Save BLAST results and GenBank Report to your computer or diskette and print out the file. The latter should look like the example below after changing font to the proportional Courier font at small size 8pt.

```
LOCUS HSPREC 5003 bp RNA PRI 12-SEP-1993
DEFINITION Human mRNA and promoter DNA for progesterone receptor.
ACCESSION X51730
NID g35651
KEYWORDS hormone receptor; progesterone receptor.
SOURCE human.
ORGANISM Homo sapiens
Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE 1 (bases 1 to 5003)
AUTHORS Kastner,P.
TITLE Direct Submission
JOURNAL Submitted (16-FEB-1990) Kastner P., LGME/CNRS - U184/INSERM, 11 rue
Humann, 67085 Strasbourg Cedex, France
```

```
REFERENCE 2 (bases 1 to 5003)
AUTHORS Kastner,P., Krust,A., Turcotte,B., Stropp,U., Tora,L.,
Gronemeyer,H. and Chambon,P.
TITLE Two distinct estrogen-regulated promoters generate transcripts
encoding the two functionally different human progesterone receptor
forms A and B
JOURNAL EMBO J. 9 (5), 1603-1614 (1990)
MEDLINE 90228361
COMMENT See also <M15716>.
Bases 1-711 were derived from genomic DNA.
FEATURES Location/Qualifiers
source 1..5003
/organism="Homo sapiens"
/db_xref="taxon:9606"
/cell_type="T47D"
/clone_lib="lambda-gt11"
/chromosome="11"
/map="q22"
misc_feature 712
/note="beginning of mRNA sequence"
CDS 1455..4256
/note="progesterone receptor (AA 1-933)"
/codon_start=1
/db_xref="PID:g35652"
/db_xref="SWISS-PROT:P06401"
/translation="MTELKAKGPRAPHVAGGPPSPEVGSPLLCRPAAGPFFGSQTSMT
LPEVSAIPIISLDGLLFPRPCQGQDPSEKTDQDQQLSDVEGAYSRAEATRGAGSSSS
PPEKDSGLLDVLDTLAPSGPGQSPSPACEVTSSWCLFGPELPPDPPAAPATQRV
LSPLMSRSGCKVGDSSGTAHAKVLPRLGSPARQLLLPASESPHWSGAPVKPSPQAAA
VEVEEDSSEESAGPLLKGPRLGGAAGGAAACPPGAAAGGVALVPKEDSRFS
APRVALVEQDAPMAPGRSPLATTVMDFIHVPILPLNHALLAARTRQLEDES YDGGAG
```

AASAFAPRRTSPCASSTPVAVGDFPDCAYPPDAEPKDDAYPLYSDFPALPKIKEEEE
 GAEASARSPRSYLAVAGANPAAFPDPFLGPPPLPPRATPSRPGEEAAVTAAPASASVSS
 ASSSGSTLECILYKAEGAPPQGGFFAPPPCKAPGASGCLLPRDGLPSTASAAAAAGAA
 PALYPALGLNGLPQLGYQAAVLKEGLPQVYPPYLNLYLRPDSEASQSPQYSFESLPQKI
 CLICGDEASGCHYVLTGCSCKVFFKRAMEGQHNYLCAGRNDCIVDKIRKNCPCARL
 RKCCQAGMVLGGRRKFKFNKVRVVRALDAVALPQLGVPNESQALSQRFTFSPGQDIQ
 LIPPLINLLMSIEPDVIYAGHDNTKPDTSSSLLTSLNQLGERQLLSVVKWSKSLPGFR
 NLHIDQITLIQYSWMSLMVFLGWRSYKHVSGQMLYFAPDLILNEQRMKESSFYSLC
 LTMWQIPQEFVKLQVSQEEFLCMKVLLLLLNTIPLEGLRSQTQFEEMRSSYIRELIKAI
 GLRQKGVVSSSRQFYQLTKLLDNLHDLVKQLHLHYCLNTFIQSRALSVEFPEMMSEVIA
 AQLPKILAGMVKPLLFHKK "

BASE COUNT 1233 a 1303 c 1240 g 1227 t
 ORIGIN

```

1 ggatccattt tataagctca aagataatta cttttcagac taagaatatt tagggtaaaa
61 agtactgttc aacatctcta ctgaggatgt tatgatgtag cacactctat aagctggagc
121 taaaggaaac tttccttaaa gtgctattta ctaaaaattg gaacacattc cttaagacaa
181 atcgaagtgt ggcacacaac atccaaactt ccatcataga tacagagggt ttaccatctc
241 ccaactccaa atttctttgt cacgctgagg ataactcaaga ggagcaggac atgttggtcg
301 cagcaggaga aacttgaag cattcacttt tatggaactc ataagggaga gaatctctta
361 tttagatcgc tccttgatagc atttattatt ttaaaagata atgtagccaa atgtcttctc
421 ctgtgttaaa tctttacaaa actgaaatct taaaatggtg acaaaaattc tacttctgat
481 agaactctatt ctttttcca attagatagg gcataattct taatttgcga acaaaaacgt
541 aatatgctta tgaggttcca tcccaaagaa cctgctattg agagtagcat tcagaataac
601 ggggtgaaat gccaaactcca gagtttcaga tcctaccggg aattggggta gggaggggct
661 ttgggcgggg cctcccctaga ggaggaggcg ttgtagaaa gctgtctggc cagtccacag
721 ctgtcactaa tcggggtgag cctgtgtgta ttgctcgtg tgggtggcat tctcaatgag
781 aactagcttc acttgtcatt tgagtgaat ctacaaccg aggcggttag tgctcccga
841 ctactgggat ctgagatctt cggagatgac tgtcgcggc agtacggagc cagcagaagt
901 ccgacccttc ctgggaatgg gctgtaccga gaggtccgac tagccccagg gttttagtga
961 gggggcagtg gaactcagcg agggactgag agcttcacag catgcacgag tttgatgcca
1021 gagaaaaagt cgggagataa aggagccgcg tgtcactaaa ttgccgtcgc agccgcagcc
1081 actcaagtgc cggacttgtg agtactctgc gtctccagtc ctccgacaga agttggagaa
1141 ctctcttgga gaactccccg agttaggaga cgagatctcc taacaattac tactttttct
1201 tgcgctcccc acttgcgctc cgctgggaca aacgacagcc acagttcccc tgacgacagg
1261 atggaggcca agggcaggag ctgaccagcg ccgcccctccc ccgcccccca cccaggaggt
1321 ggagatcctc cgggtccagc acattcaaca cccactttct cctcccctctg cccctatatt
1381 cccgaaaccc cctcctcctt cccttttccc tctcctctgg agacggggga ggagaaaagg
1441 ggagtccagt cgtcatgact gagctgaagg caaagggtcc cggggtccc cagtgggcgg
1501 gcgcccgcgc ctcccccgag gtcggatccc cactgctgtg tcgcccagcc gcaggtccgt
1561 tcccggggag ccagacctcg gacacctgc ctgaagtttc ggccatacct atctccctgg
1621 acgggctact cttcctcctg cctgcccagg gacaggacc ctccgacgaa aagacgcagg...

```

8. So now, in the example, I know my cDNA clone is part of progesterone receptor mRNA, probably human.

Note: It is important to remember that my cDNA is related to the GenBank entry but may not be identical to it.

9. I can determine the whether the cDNA sequence is on the sense or antisense strand by looking at the position numbers of the subject sequence in the alignment.

```

>emb|X06624|OCPRR3 Human progesterone receptor mRNA 3'-flank
Length = 3566
Score = 109 bits (55), Expect = 3e-23
Identities = 62/63 (98%), Positives = 62/63 (98%), Gaps = 1/63 (1%)

Query:      1 attattgaagtaagctatgtcttaccatactatttcataccatttaagtgaggatatt 60
             |||
Sbjct:     273 attattgaagtaagctatgtcttaccatactatttcata-ccatttaagtgaggatatt 331

```

Since the numbers are ascending along with the Query numbers, my query is sense strand, which is the standard entry for mRNA sequences in GenBank.

FOR YOUR TWO SEQUENCE ANALYSES:

1. Do the analyses at NCBI described above and save and print the top hit alignments and their GenBank entries.
2. Answer the following questions about the sequences you analyze in your notebooks:

Q1: The cDNA is _____ (sense or antisense) from _____ mRNA.

Q2: IS YOUR GENE PRODUCT CONSERVED ACROSS SPECIES? Can you tell which species your sequence is from? Did you get hits from distantly related species or not? (*Note: GenBank has more human and mouse sequences in it than those of other mammals.*)

Q3: WHAT IS THE FUNCTION OF THE GENE PRODUCT? Look at the protein/gene name and titles of the references in the GenBank report of the top hit.

Q4: Is the Query sequence within the coding sequence? If so, what amino acids does it encode? Can translate by hand or using a translation tool on the web.

Q5: WHAT TISSUES IS YOUR GENE PRODUCT EXPRESSED IN? (You can go to the human UniGene entry for PGR (gene name) to see the EST Profile. There are also GeoProfile entries of microarray datasets that can tell you a lot about that gene's expression and regulation)

GENE EXPRESSION Lab 9 ning 11/5/10

IN VITRO TRANSCRIPTION OF ANTISENSE cRNA PROBES

[Be RNase free...and use aerosol barrier tips! See in vitro transcription (as in Biotin-16-UTP (Roche) protocol) and Maxiscript (Ambion) protocols]

TODAY each student will make an antisense cRNA probe for aromatase mRNA and one to an “unknown” RNA for hybridizing to the Northern blots. The former will use the purified PCR band in low melt gel as a DNA template. The lab boss will give out a linearized plasmid containing complementary DNAs (cDNA's, synthetic copies of fragments of mRNAs) for the latter.

There are 3 common types of nucleotide probes: DNA oligonucleotides (ss), cDNA (ds) and cRNA (ss). For many applications, cRNA probes are superior over:

1. end-labelled oligonucleotide probes because they are:
 - a. longer (and therefore carry more label and have higher hybridization specificity)
 - b. uniformly labelled throughout (so they carry more label)
2. nick-translated or random-primed cDNA probes, because they only have the desired probe strand, not the other "sense" strand that increases background.

In addition, the binding of RNA:RNA hybrids is stronger than that of DNA:DNA hybrids.

To make cRNA probes, a DNA template is needed with a bacteriophage promoter site 5' to the cDNA. For antisense cRNA that will bind mRNA, the promoter is on the 5' side of the antisense strand. For transcribing sense cRNA (*Q: What is that useful for?*) the promoter sequence would be 5' to the sense strand. DNA templates can be PCR products or circular plasmid bearing cDNAs that are restricted or cut at a specific site with a restriction enzyme where the transcript will terminate.

For in vitro transcription, the DNA template and ribonucleotides are combined with a bacteriophage RNA polymerase (either T7 or T3 or SP6, in order of efficiency). The polymerase enzyme binds a specific site (its promoter) on the plasmid and transcribes (makes RNA) using the DNA as a template.

REMEMBER: the cDNAs are synthetic cloned fragments of the mRNAs
The antisense cRNAs synthesized today will be labeled with biotin so they will be probes (detectable reagents) for identifying its homologous mRNA in the tissue RNA samples. After hybridizing the probe to the RNA on the Northern blot and washing the blot, specifically bound probe will be detected with either streptavidin (which binds biotin). conjugated to alkaline phosphatase enzyme, which will cleave the chemiluminescent detection reagents to yield a low intensity light signal that is detected by X-ray film.

*Be clean – clean and protect apparatus from RNase contamination- work on clean
diapers! Use all DEPC reagents, gloved hands and aerosol barrier tips.*

I. START THIS FIRST! In vitro Transcription.

Each student should set up two in vitro transcription reactions for biotin-labeling antisense cRNA probes: one for aromatase mRNA and one for the unknown RNA.

1. Thaw components at room temperature (RT) then store on ice.

EXCEPTION: RNasin and RNA Polymerase, like all enzymes, stay at -20°C always! and

Melt the aromatase template in low melt gel at 70°C for 10 min. Then put the tube in the 37°C block for > 5min. You will pipet the template from the tube at that location. Keep the other template (in aqueous phase) on ice.

2. For each antisense cRNA synthesis, add components from kit, in order, to a 0.5 ml tube at RT.

_____ ul DEPC-H₂O to make final reaction volume 20 ul.

4 ul 5X Transcription Buffer (Promega)

1 ul RNasin

2ul 100 mM DTT

2ul 10 mM rATP, rCTP, rGTP, and 3.5 mM Biotin-16-UTP

1 ul DNA template (aqueous) or 3 ul template in low melt gel

1 ul T7 RNA Polymerase

3. Mix by pipetting up and down...gently! No bubbles.

4. Flash spin in a microfuge

5. Incubate 37°C for 1h

6. Add 1 ul RNase-free DNase

7. Incubate 15 min at 37°C

8. Place two 1 ul aliquots of each reaction into labeled tubes for the next lab.

9. Store all of the RNAs at -80°C.

GENE EXPRESSION Lab 10 ning 11/12/10

Assess quality of in vitro transcription products.

A. Analysis of in vitro transcribed probes on an acrylamide gel ("probe test gel", recipe in Appendix).

1. A group of four students should make one 5% polyacrylamide/ 8 M urea gel, RNase-free, in a vertical slab gel apparatus (see Appendix for gel recipe). Add comb and allow to polymerize 20 min. **NOTE: unpolymerized acrylamide and TEMED are TOXIC!!**
2. Set up gel rig with 1.5 liters of 1X TBE in DEPC water.
3. Add 10 ul RNA Load dye containing bromphenol blue and xylene cyanol dyes to each 1 ul aliquot from the in vitro transcription reactions and mix. Ask if RNA Century markers from Ambion are available. Heat all samples for the gel at 70°C for 10 min. then put on ice. Then load on the gel after flushing the wells free of urea that leaches into them from the gel.
4. Run at 35 mamps for 30 min.
5. Stain the gel with EtBr (agitate in 100 mls of 2.5 ug/ml EtBr in DEPC H₂O for 5 to 10 min , then destain similarly in DEPC H₂O) and photograph on UV box.

QUESTIONS: 1. Are both probes of high quality? (of expected size and present in high concentration?)

2. If you used Century markers, based on the masses of those bands, estimate the RNA concentration of your probe solutions.

3. Compete with your Northern blot partner....which aromatase and unknown probes look best and should be used for blot hybridization?

B. Dot blot of serial dilutions of probe: (After 1 h transcription)

1. Make 4 serial 1:9 dilutions of each probe: Pipet 1 ul undiluted probe into a tube with 9 ul TE and mix. This is a 1:10 dilution. Repeat the dilution 3 times; so have 1:10, 1:100, 1:1000 and 1:10,000 dilutions. (...or 10⁻¹, 10⁻², 10⁻³, and 10⁻⁴ dilutions)
2. On one 3 by 10 cm strips of Brightstar nylon membrane, mark membrane with sharp pencil or black Sharpie pen "-1", "-2", "-3", "-4" at 2 cm intervals across the width of the blot. Label rows of the dot blot "aromatase" and "unknown". Add initials and date.
3. Dot 1 ul of appropriate probe dilutions under labels. UV Cross-link RNA to membrane as you did for the Northern blot. Air dry and store in a clean place between paper towels. (You'll develop the dot blot along with the Northern blot on 11/19/10.)

GENE EXPRESSION Lecture/Lab 11 ning 11/18 & 19/10

Thursday 11/18/10 in class

Northern blot Prehybridization & Hybridization

Each pair of students will hybridize the top half of the northern blot with aromatase probe and the bottom with the “unknown” probe. Since the nylon membrane likes to bind things, background sites are blocked (bound) with non-specific DNA and protein.

Usually, sheared salmon sperm DNA is used in prehybe to block these sites.

1. Warm the Ambion Ultrahybe hybridization solution to 55oC. Swirl to dissolve precipitates. Cut the shared Northern blot to separate top and bottom halves so that student 1 and student 2 samples on the top can be probed for aromatase mRNA and the bottom half of the blot can be probed with the unknown probe.
2. Wash two hybridization bottles with RNaseZAP. Insert one dry blot into each bottle so the RNA side is toward the center. Wet the blots with 2X SSC. Pour 2X SSC out.
3. Add 10 to 20 ml Hybridization solution to each and seal bottle screw caps.
4. Incubate 55oC for 30 min in the hybridization oven with bottles rotating.
5. From the estimated concentrations of probes, calculate the volume of each probe to add to 10 or 20 ml hybridization buffer so that the concentration is 0.1 nM.
6. Heat the cRNA probes from the last lab at 94oC for 10 min - chill on ice. Add ____ ul (from #5) of heat denatured in vitro transcribed probe to the appropriate hybridization solution bottle.
7. Incubate 55oC overnight with hybridization bottles rotating.

DO THE FOLLOWING IN YOUR NOTEBOOK:

1. Based on your blot photo and what you know, **draw your expected hybridization results** in your notebook. Use the 28S & 18S rRNA positions as markers. For the mouse, 28S rRNA is 4718 bases and 18S rRNA is 1847 bases.

And answer QUESTIONS:

2. **How long are the aromatase and “unknown” mRNAs?**
3. **In one of the RNA samples from tissue, how much RNA is in each rRNA band?**

LAB 11 - Friday 11/20/09

Northern Blot Washing and Development

[See Biotin luminescent detection kit (Roche) and NorthernMax (Ambion) protocols]

After overnight hybridization, probe is maximally bound to specific sequences. It is also present on some non-specific sites. By reducing [salt], mainly in the form of SSC, hybridizations are tested for stringency. Usually temperature is increased as well so that only specific probe-binding for the target of interest remains after washing.

KEEP BLOTS WET DURING THESE PROCEDURES or you'll have a lot of artifacts.

Wash blots with vigorous shaking and seal the tupperware lids during washes so solutions don't spill onto the shakers!!! When binding reagents use gentle agitation.

1. Open the hybridization bottles and discard hybridization solution down the drain.
2. Move the blots to a clean tupperware container with a sealing lid. Can wash several blots in the same container if they are distinguishable from each other (sometimes the labeling fades overnight with this hybridization solution).
3. Wash the blot in 100 mls of 42°C 2x SSC [or "low stringency wash" from Ambion] with hard shaking for 15 min at 42°C.
4. Discard wash and rewash with 100 mls of 0.1X SSC at 42°C.

REST OF STEPS AT ROOM TEMPERATURE:

5. Wash in 50 mls Washing Buffer [Maleic acid buffer (1X = 0.1 M maleic acid, 0.15 M NaCl, pH 7.5) with 0.3% Tween-20] at RT for 5 min.

6. Repeat #5 once. Add in the probe dot blot from the previous lab at this step.

7. Transfer to a clean, freshly washed Tupperware container or tip lid and incubate the membranes for 30 min in 20 ml Block solution (1% (w/v) blocking reagent in maleic acid buffer without Tween). During gentle shaking, membranes should move independently from each other.

8. Discard Block and incubate in Streptavidin-Alkaline Phosphatase solution (1:5,000 diluted in block solution) for 30 min. Use a minimal volume (20 ml) in a small clean container like a yellow tip box lid. During gentle shaking, membranes should move independently from each other.

9. Discard the antibody solution and wash twice in Washing Buffer for 15 min each time.

10. Discard wash and equilibrate membrane in Detection Buffer [0.1 M Tris/HCl(pH9.5), 0.1 M NaCl] for 2 min.

11. Pipet 1 ml of room temperature CDP-Star Detection Reagent diluted 1:100 in detection buffer onto each blot and cover with plastic wrap. Incubate 5 min, then pour off excess reagent.

12. Wrap blot in saran wrap. To keep the blot wet and the film dry, double fold the plastic wrap and tuck all edges under the blot. As always, RNA side up!

13. Place in cassette.

14. Go to the dark room and lay a piece of X-ray film on the blot. Bend the lower right corner of the film to create a dark line for film orientation. Close the cassette. Make sure you fold the flaps correctly so it is light tight!

15. Place cassette at 37°C or room temperature. Develop the film in 10 min in the developer machine in the dark room. Label films with exposure date, time and index to your notebook. Align the film with the blot and mark the positions of the wells, 28S & 18S rRNA migration distances on the film. Identify and label lanes. If desired, place unexposed X-ray film on the blot and expose longer...overnight?.

Review your predicted Northern blot results from the previous class session. It is imperative that you know what you are looking for in order to assess and optimize the blot exposures...

GENE EXPRESSION LECTURE 12/6/10

REVIEW RESULTS

&

NORTHERN BLOT ANALYSIS:

1. Qualitatively assess the blot results:

- a. Is exposure optimal?

- b. How many bands are evident?

- c. Are they in the expected tissues?

- d. What are the sizes of the hybridizing bands with respect to the rRNAs?

- e. For one band, empirically determine the length of the mRNA hybridized. To do this measure migration distances for the band on the Xray film, and for 28S (4800 bases) and 18S (1800 bases) rRNAs on the EtBr stained gel photo using UV ruler as a guide. On graph paper, you can plot log base length against migration distance for the rRNAs. Find log bases from the plot, using the migration distance of the band of interest. Find antilog to get number of bases. If RNA markers are used, you can use those instead of or in addition to the 18 and 28S rRNAs.

2. Quantitate hybridizing bands

- a. Can use densitometry on the X-ray film. Need a good scanner and analytical software such as BioImage IQ.

- b. Many machines are being developed for direct scanning of blots. These are very powerful because they avoid the limitations of film and generate electronic data files for easy storage and publication.

3. CLEAN UP THE LAB and REMOVE PERSONAL ITEMS.

APPENDIX

Use of Micropipettors

1. Choose the correct pipet. For volumes:
 - 1-20 ul P20
 - 20-200 ul P200
 - 200 - 1000 ul P1000
 2. Set the desired volume by holding the pipetman in one hand and turning the volume adjustment knob until the correct volume shows on the indicator. For best precision, always approach the desired volume by dialing downward (at least one-third revolution) from a larger volume setting.
 3. Attach a new tip to the shaft of the pipet. Press tip on firmly to ensure airtight seal. Choose the correct tip.
 - P20 yellow tip
 - P200 yellow tip
 - P1000 blue tip
 4. Depress plunger to first positive stop. Hold pipetman vertically and immerse disposable tip into sample liquid 2mm.
 5. Allow the push button to return slowly to the up position. Never permit it to snap up.
 6. Wait 1 or 2 seconds to ensure that the full volume of the sample is drawn into the tip.
 7. Withdraw tip from the sample liquid. Wipe the sides of the tip on the sides of tube to remove any remaining liquid.
 8. To dispense the sample, place the tip end against the side wall of the receiving vessel and depress the plunger slowly to the first stop. Then depress the plunger to the second stop to expel any residual liquid in the tip.
 9. With the plunger fully depressed, withdraw pipetman from the vessel. Then allow the plunger to return to the top position.
 10. Discard tip by depressing the tip ejector button. A fresh tip should be used for each sample.
-

Terribly Difficult Calculations

1. Molar solutions

1 M (mole per liter) means the solution has 1 molecular weight mass (g) per volume (liter) of soln.

A mole is a number of molecules:

6.022×10^{23} , Avogadro's number

To make 500 mls of 0.5 M NaCl (NaCl is 58.55 g/mole) you need (0.5 liters)

(0.5 mole) = 0.25 mole

liter)

$0.25 \text{ mole} * 58.55 \text{ g/mole} = 14.6 \text{ g}$

So: Add 14.6 g NaCl powder and bring final volume to 500 ml with H₂O.

2. We typically work with concentrated stock solutions. For example, our Tris/acetate/EDTA (TAE) is made as a 50X stock. We run gels in 1000 mls of 1X TAE.

The way I do DILUTION PROBLEMS is:

$[\text{Stock}] * y = [\text{Desired}] * \text{Desired volume}$; where y is the volume of stock. To find Y needed to make 1 li of 1X TAE from a 50X stock:

$50X * y = 1X * 1000 \text{ ml}$

$y = 1X/50X * 1000 \text{ ml} = 20 \text{ ml}$

So add 20 ml 50X TAE to a 1 liter graduated cylinder. Bring volume to 1 li w/ dH₂O.

3. Note: Dilutions are applicable to problems of pipetting very small amounts. If you want to add 0.2 ul, dilute the material 1:9 and pipet 2 ul with a P-20.

4. Percentage solutions should have a (v/v) or (w/v) or (w/w) following.

a. (v/v) relates volume to volume, indicating both components are liquids: e.g. 100 mls of 75% (v/v) EtOH is made with 75 mls EtOH + 25 ml H₂O

b. (w/v) indicates solid to liquid ratio: e.g., 10 mls of 10% (w/v) ammonium persulfate (APS) is made w/ 1 g of APS to 10 ml final volume with water.

c. (w/w) is rare, indicating a weight to weight relationship. To make 10 mls of a 10% (w/w) APS soln, you could weigh 1 g APS on a scale and then add water until solution weight is 10 g. (That would be 9 g = 9 mls since density of H₂O is 1 g/ml).

5. Of course, these calculations can be combined. For example, to make 500 mls of 0.5 M NaCl in 1X TAE,

Combine 14.6 g NaCl with 10 mls 50X TAE. Bring volume to 500 mls with H₂O.

Easy!

ACRYLAMIDE “PROBE TEST” GEL

-

(short, fat sequencing gel)

5% acrylamide/urea gel

25.5g urea
19.5ml H₂O
12ml 5X TBE
7.5ml 40% (w/v) acrylamide (19:1 acryl:bis)
60ml final vol.

Heat to 37°C to dissolve urea

Cool to below RT

Filter (~optional)

Add 400 ul 10% APS (less than 1 week old, make 1ml)

50 ul TEMED

Clean gel plates with soap, rinse with H₂O extensively, then wipe with EtOH and Kimwipes. Set up plates as instructed.

Pour into 1.5mm thick vertical gel slab. Add comb.

Bubbles = Bad

Should polymerize in 15-20 min.

Rinse wells with 1X TBE immediately after pulling the comb and just prior to loading samples.

Prepare samples in 80% (v/v) formamide loading dye

Heat 68°C 5 min for RNA. For DNA, 94°C 5 min.

Run at 25 to 35 mamps.

Dye Migration Related to bases in a denaturing gel (d) or base pairs in a non-denaturing gel (n):

<u>Acrylamide %</u>	<u>Bromphenol Blue</u>	<u>Xylene</u>
<u>Cyanol</u>		
3.5% n	100	-
5% n	65	-
5% d	35	130
6% d	26	106
8% n	45	-
8% d	19	75
10% d	12	55
12% n	20	-
20% n	12	-

e.g., in a 5% acylamide + urea gel (denaturing), Bromphenol Blue comigrates with 35 base long nucleic acids.

Making Agarose Gels for DNA

For one gel, make 1 li of 1X TAE from the 50X stock in a 1 liter graduated cylinder and mix well by inversion. Use 100 ml to make the gel and the rest to run the gel.

1%	0.8 % Low melting point agarose gel
1g Agarose (regular) agarose	0.8 g Low melting point
100ml 1X TAE	100ml 1X TAE
10ul 10mg/ml EtBr	10 ul 10mg/ml EtBr

Put components in a 500 or 300 ml beaker, cover with Saran Wrap, heat in microwave until solution boils 3 times. Allow to cool to 60oC, pour gel into mold as instructed.

NOTE: Low melting agarose gels are very pure chemically and are used to prepare or purify DNA. However, they are hard to see and work with...slippery, easy to break and generally annoying. DNA bands will be less defined on them, as well.