# Tobacco Induced Mutation: Teacher Instructions 

Summary: Students expose bacteria to 4 different concentrations of tobacco extract and observe the mutagenic effect on the bacteria. (The bacteria change from red to white when they mutate). Mutations are an important precursor to cancer.

Objectives: Students will leam how to plate bacteria.
Students will leam how to collect \& a nalyze data.
Students will observe the mutagenic properties of tobacco.
Students will observe a dose/response relationship.
Time: Preparation - Make Agar Petri Plates: 2 hours
Make Stock Plates: 10 minutes
Make Tobacco Solution:30 minutes
Experiment - Perform the Experiment: 1 hour
Incubate Bacteria: 24-48 hours (doesn't require class time)
Data Collection \& Analysis: 1-3 hours (depending on extent of a nalysis)
Materials:

| Material | Vendor and Catalog Number |
| :--- | :--- |
| Petri Dishes (1-2 Plates per student) | VWR 100 mm sterile 25384-071C a se <br> of $500 \$ 96.30$ |
| Wire Loops (one for each student) |  |
| Tryptone | Difc $, 211703,100 \mathrm{~g}$ |
| Yeast Extract | Difco $, 212740,100 \mathrm{~g}$ |
| Cigarettes (2 per group) |  |
| NaCl (Sodium Chloride) |  |
| UV la mp |  |
| Beaker (one per group) |  |
| Stir sticks |  |
| Hot plate or mic rowave |  |


| Test tube racks which fit 14 ml tubes <br> (1 per group) |  |
| :--- | :--- |
| Cheese Cloth or other Filter Paper <br> (1 per group) |  |
| 3 500 ml Bottles |  |
| Bunsen Bumers (3-4) |  |
| Graduated Cylinder | VWR 60819-761, case of 500, 111.37 |
| 1 ml Size Pla stic Tra nsfer Pipettes (4 <br> per group) | VWR 14670-345 500 for \$24.78 |
| 14 ml Plastic Test Tubes (5 per <br> group) | Difco, 214010, 454g |
| Agar |  |
| Autoclave or pressure cooker |  |
| Serratia Marcescens |  |
| Distilled Water |  |
| Marking Pens (1 per student) |  |

## Methods:

TEACHER PREPARATION
Make the Agar Petri (LB) Plates and liquid bacterial media (food); allow minimum two days to complete the preparations

- LB- Plates:

Mix the following ingredients with 1 L of distilled water ( 20 ml media per plate, makes about 35 plates):

## 10 g NaCl

10g tryptone
$5 g$ yeast extract
20 g agar
Autoclave the solution ( $121^{\circ} \mathrm{C}, 30$ minutes); wa it for the solution to cool (should be cool enough to handle but not cold) and pour carefully into the Petri dishes avoiding bubblesas much as possible. Allow gel to solidify at room temperature ovemight ( $\mathrm{O} / \mathrm{N}$ ). BB-Platescan be stored in a refrigerator for up to 2 months or at room temperature for a couple of days.

Tips:

1) use a 2 L glass Erlenmeyer flask for the mixture to prevent bubbling over of the mixture during heating, cover the top of the Erlenmeyer with foil before autoc la ving.
2) add all ingredients except agar initially; once all the other ingredients are in solution, add the agar. The agar will not go into solution at room temperature but this is okay as the agar will go into solution during a utoc laving.
3) Whether inoculated with bacteria or not, store the plates upside down to a void condensation buildup on the surface of the agar.

- Liquid LB-Media:

Mix the following ingredients with 1 L of distilled water.
10 g NaCl
10g tryptone
$5 g$ yeast extract
Transfer solution to $2-500 \mathrm{ml}$ glass bottles and a utocla ve ( $121^{\circ} \mathrm{C}, 30$ minutes). This solution should be stored in the bottles and can be stored at room temperature indefinitely.

## Prepare bacteria and tobacco solution for the experiment

- Serratia Marcescens: S. Marcescens is the strain of bacteria utilized in this a ctivity and is non-infectious to humans. S. Marcescens produces a red pigment (i.e. colonies are red) when grown at $30^{\circ} \mathrm{C}$. When grown at cooler or wa mer temperatures, this red pigment is not produced and colonies are white. Mutations in the bacteria will produce white colonies at $30^{\circ} \mathrm{C}$. Thus, it is important to grow the bacteria at $30^{\circ} \mathrm{C}$ when testing for the a bility of the ciga rette extract to produce mutations.
- Generate Stock Plates: After obtaining S. Marcescens, the tea cher should make some stock plates which will serve as a reservoir of bacteria for activities with students. Stock plates can be generated by:

1) inoculating 3-4 LB-plates with S. Marcescens
2) incubating the plates ovemight at $30^{\circ} \mathrm{C}$. These plates remain viable and can be stored up to several months in the refrigerator.
Generate 3-4 "student stock plates" from stock plates of Serratia marcescens for the students to use in order to prevent contamination of the refrigerated stock plates: Prior to a ny activity with students, 3-4 student stock plates should be generated by:
3) fla ming wire loop
4) removing a small amount of bacteria from the refrigerated stock LBplates using the wire loop.
5) Inoculating 3-4 new $\operatorname{B}$ plates with the bacteria on the wire loop, incubate $24-48 \mathrm{hr}$. at $30^{\circ} \mathrm{C}$.

- *Make Tobacco Solution:

1) Remove the tobacco from 2 cigarettes and place in 200 ml of distilled water in a beaker.
2) Heat the water for $10-15$ minutes with constant stiming .
3) Filter the leavesout of the extract using cheese cloth or other filter paper.
4) Pour into a bottle forstorage
*This step can be performed by the teacher or by the students depending on time constraints.

## DAY 1. PERFORMING THE EXPERIMENT: 1 hour

- Place the 3-4 "student stock plates" a round the room.
- Divide the students into groups of $4-5$ a nd distribute 6 LB media plates to each group and a wire loop to each student.
- Have students label 5 tubes: 1:1, 1:100, 1:1000, 1:10,000, bacteria
- Have students make dilutions of the cigarette solution using graduated pipettes according to the scheme below.

1:1 - no dilution, place 0.1 ml of cigarette solution into the 14 ml tube labeled 1:1
1:100 - place 0.1 ml cigarette solution +10 ml liquid BB media into the 14 ml tube labeled $1: 100$, shake solution to mix
1:1000 - using a new pipet, place 0.1 ml of the 1:100 ml cigarette solution +10 ml LB media into the 14 ml tube labeled 1:1000, shake solution to mix
1:10,000 - using a new pipet place 0.1 ml of the 1:1000 cigarette solution +10 ml LB media into the 14 ml tube labeled 1:1000, shake solution to mix
Bacteria - The students should save this tube for later. Further instructions for this tube will be given later in this protocol.

- Have one student in each group prepare an LB-plate with one of the above cigarette solutions. The teachermay want to point out the position of 0.1 ml mark on the graduated pipet. To prepare the plates, have students:

1) label the bottom of the plate with theirname, date, and NT(no treatment), UV (UV light Positive control), 1:100, 1:1000, or 10,000
2) add 0.1 ml of dilution solution to properly labeled LB-plate
3) bend wire loop into an Lshape
4) flame loop
5) carefully spread 0.1 ml of dilution solution a round the agar with the wire loop bent in the shape of an $L$
6) Flame the loop again to sterilize for the next step.

## Make sure the students remember to use a different pipet for each of the different dilution solutions to prevent contamination!!

- Let the plates sit right-side up (large lid on top) at room temperature to allow the ciga rette solutions to sink into the surface of the Agar Petri Plates (approximately $20-30$ minutes, can be left ovemight due to time contraints if need be).


## Treatment Conditions

| Tobacco Solution | Controls |
| :--- | :--- |
| $1: 1$ | No treatment $=$ NT <br> (negative) |
| $1: 100$ | UV light (positive) |
| $1: 1000$ |  |
| $1: 10,000$ |  |

1) While the cigarette solutions are allowed to sink into the LB-plates students should:
2) Using a graduated cylinder, place 10 ml liquid B -media in the 14 ml plastic tube previously labeled-bacteria.
3) flame a wire loop
4) once the loop has cooled, remove a very small a mount of bacteria from the "student stock plates" situated a round the room.
5) The bacteria should then be placed in the 10 ml of B liquid media in the 14 ml plastic tube to dilute the bacteria. The students should "stir" the loop around in the LB media and watch to make sure the "chunk" of bacteria comes off the loop.
6) Students should then GENILY TURN THE TUBE UPSIDE DOWN 4-5 times to mix the bacteria.

- Once the cigarette solutions have been allowed to sink into the surface of the agareach student should inoculate the labeled plates with bacteria.

1) unbend wire loop
2) flame loop and wait for the loop to cool
3) dip the wire loop into the tube labeled Bacteria
4) Use the wire loop to plate the bacteria on the appropriate LB-plate using the plating pattem included in the "Student Guide to Aseptic Technique". Reflame the loop to sterilize forstorage.

## Students should make only 1 pass with their loop, while plating, otherwise, bacteria end up all over the plate and no colonies will be seen!

5) The UV light positive control plate and NTnegative control plates should be plated identically to the treatment plates but without cigarette solution.
6) The teacher should expose the UV light plate to UV light by:

- removing the Petri dish lid
- placing the open plate in front of a UV light for 60 seconds.

7) Incubate plates at $30^{\circ} \mathrm{C}$ for 48 hr .

Note: Plating bacteria should be done utilizing aseptic (sterile) technique. A description of plating bacteria utilizing a septic technique is included in the "Guide to Plating Bacteria Using Sterile Technique". The instructor may want to set aside a class period to allow the students to practice working with bacteria.

## Day 2: BACTERIA INCUBATING

## Day 3-4: DATA COШECTION AND ANALYSIS: 2-3 hr.

Collecting the data:

- Each student should obta in his/her plate and begin to count the white colonies.
- The number of white colonies for each dilution should then be recorded in a laboratory notebook or on the data sheet provided.
- If no colonies are seen on the students plate and the bacteria grow in a lawn, data can be collected asan estimation of the percentage of white bacteria on the plate. Percentage can be easily determined by utilizing the grid included in this packet.

1) Students should place their previously inncoulated petri plate down onto the grid, lining up the outline of the circle with the outline of the bottom of their plate.
2) The students should then estimate the percentage of white bacteria, red bacteria, or empty of each of the 32 small squares
and record the percentages in the appropriate column next to the grid circle.
3) The total percentages of each colorcan then be calculated by adding up the percentagesfrom all 32 small squares and dividing the total by 32 .

## Analyzing the Data : Disc ussion Points

- Did the students see a ny trends in their group? (e.g. The number of mutations increase as the tobacco solution concentration increases)
- Why are mutations important?
- Did any groups have mutations at the 1:10,000 dilution? If so, discuss how small that a mount is.

Statistical Analysis (optional): The instructor may collect the data for the whole class and post it such that each student can a nalyze all the data. An overhead for the class data sheet is included in this packet.

- Once all of the data has been collected the average and standard deviation for the number of white colonies for each treatment condition can be calculated.
- T-testscan then be performed to determine statistic al signific ance of the results. The following comparisons might prove interesting:

NTX 1:1 (no treatment negative control compared to 1:1 dilution)
NTX 1:100
NTX 1:1000
NTX 1:10,000
NTX UV light

- Additionally, the results can be graphed as number of colonies vs. dilution of ciga rette solution on graph paper such asthat included in this packet.


# Tobacco Induced Mutations: Student Instructions 

## Materials:

6 Petri Dishes
Wire Loops (one for each group
member)
Cigarettes (2)
Beaker
Stir stick
Hot plate or mic rowave LB-Plates (bacteria food)
LB liquid media (bacteria food)

Test tube racks which fit 14 ml tubes Cheese cloth or other filter paper<br>1500 ml bottle<br>Bunsen Bumer<br>Graduated Cylinder<br>4 1ml size plastic pipettes<br>514 ml plastic test tubes<br>Serratia marcescens<br>Distilled Water<br>Marking Pens

## Methods:

## Day 1: TREATTHE BACTERIA

Make Tobacco Solution:

- Remove the tobacco from 2 cigarettes and place in 200 ml of distilled waterin a beaker.
- Heat the water for 10-15 minutes with consta nt stiming .
- Filter the leaves out of the extract using cheese cloth or other filter paper.

Performing the experiment:

- Obta in 6 LB media plates and 5 wire loops for your group.
- Label 5 tubes: 1:1, 1:100, 1:1000, 1:10,000, bacteria
- Make dilutions of the ciga rette solution using graduated pipettes according to the scheme below. Make a check mark in the box beside the directionsfor each dilution as you make them.

1:1 - no dilution, place 0.1 ml of ciga rette solution into the 14 ml tube labeled 1:1

- place 0.1 ml cigarette solution +10 ml liquid B media into the 14 ml tube labeled 1:100, shake solution to mix - using a new pipet, place 0.1 ml of the 1:100 cigarette solution +10 ml LB media into the 14 ml tube labeled 1:1000, shake the solution to mix.
1:10,000 - using a new pipet, place 0.1 ml of the $1: 1000 \mathrm{ml}$ ciga rette solution +10 ml LB media into the 14 ml tube labeled $1: 1000$, shake solution to mix
Bacteria - Save this tube for later. Further instructions for this tube will be given later in this protocol.


## Treatment Conditions

| Ciga rette Solution | Controls |
| :--- | :--- |
| $1: 1$ | No treatment $=$ NT <br> (negative) |
| $1: 100$ | UV light (positive) |
| $1: 1000$ |  |
| $1: 10,000$ |  |

Each student in your group should prepare an BB-plate with one of the above cigarette solutions. To prepare the plate with cigarette solution:

1) label the bottom of each plate with your name \& date
2) label one plate with: NT(no treatment), UV (UV light positive control), 1:1, 1:100, 1:1000, or 1:10,000.
3) add 0.1 ml of diluted cigarette solution to the propeny la beled LB-plate using a pipette
4) bend wire loop into an Lshape
5) flame loop
6) ca refully spread 0.1 ml of dilution solution a round the agar with the wire loop.

## Remember to use a different pipet for each of the different cigarette solutions to prevent contamination!!

- Let the plates sit right-side up (large lid on top) at room temperature to allow the cigarette solutions to sink into the surface of the LB-Plates (approximately 15-20 minutes to ovemight).

While the cigarette solutions are sinking into the LB-plates:

1) Get the 14 ml tube labeled "bacteria."
2) Using a graduated cylinder, place 10 ml liquid BB -media tube.
3) Flame a wire loop
4) Once the loop is cool, remove a very small a mount of bacteria from the "student stock plates" situated a round the room.
5) Place bacteria in the 14 ml plastic tube to dilute the bacteria. "Stir" the loop around in the LB media and watch to make sure the "chunk" of bacteria comes off the loop.
6) GENTLY TURN THE TUBE UPSIDE DOWN $\mathbf{4 - 5}$ times to mix the bacteria.

Inoculate all 6 plates with bacteria:

1) unbend wire loop
2) flame loop and wait for the loop to cool
3) dip the wire loop into the tube labeled "bacteria"
4) Use the wire loop to plate the bacteria onto the appropriate LBplate using the plating pattem included in this packet. Reflame loop aftereach plating.

## Make only 1 pass with your loop, while plating, otherwise, bacteria end up all over the plate and no colonies will be seen!

5) Take your UV light plate to the teacher for exposure to UV light.

Note: Plating bacteria should be done utilizing aseptic (sterile) technique. A description of plating bacteria utilizing a septic technique is included in the "Guide to Plating Bacteria".

## Day 2: BACTERIA INCUBATING

## Day 3: DATA COШFCTION AND ANALYSIS

Collecting the data:

- Obta in your plate and count the white colonies.
- Record the number of white colonies foreach dilution on the data sheet provided.


## Analyzing the Data:

- Share yourdata with the teacher and collect the data from the other groups from the teacher.
- Calculate the average and standard deviation forthe number of white colonies for each dilution of cigarette solution from the class data sheet.
- Perform T-tests for the following comparisons:

NTX 1:1 (no treatment negative control compared to 1:1 dilution) NTX 1:100
NTX 1:1000
NTX 1:10,000
NTX UV light

- Graph number of colonies vs. dilution of cigarette solution.
- Label Axis
- Graph a verages from the class data for each dilution solution
- Paste the results of the a nalysis in your laboratory notebook.


# Tobacco Induced Mutations: Student Data Sheet 

Names of<br>Group

Members: $\qquad$
Date:

|  | Treatment Cond itions |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  | Controls |  |  | Dilution of Ciga rette Solution |  |  |
|  | No <br> Treatmen <br> t (NT- <br> negative <br> control) | UV light <br> (positive <br> control) | $1: 1$ | $1: 100$ | $1: 1000$ | $1: 10000$ |
|  |  |  |  |  |  |  |
| White (\# of <br> colonies) |  |  |  |  |  |  |

## Tobacco Induced Mutations: Class Data Sheet

|  | Treatment C ond itions |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  | Controls |  | Dilution of Ciga rette Solution |  |  |  |
|  | No <br> Treatmen <br> t (NT- <br> negative <br> control) | UV light <br> (positive <br> control) | $1: 1$ | $1: 100$ | $1: 1000$ | $1: 10000$ |
| Group 1 |  |  |  |  |  |  |
| Group 2 |  |  |  |  |  |  |
| Group 3 |  |  |  |  |  |  |
| Group 4 |  |  |  |  |  |  |
| Group 5 |  |  |  |  |  |  |
| Group 6 |  |  |  |  |  |  |
| Group 7 |  |  |  |  |  |  |
| Group 8 |  |  |  |  |  |  |
| Group 9 |  |  |  |  |  |  |
| Group 10 |  |  |  |  |  |  |
| Group 11 |  |  |  |  |  |  |
| Group 12 |  |  |  |  |  |  |
| Average |  |  |  |  |  |  |
| Standard <br> Deviation |  |  |  |  |  |  |

# Tobacco Induced Mutations: 

## Statistical Analysis

|  | Compa risons |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- |
|  | NTX UV | NTX 1:1 | NTX 1:100 | NTX 1:1000 | NTX 1:10000 |
| P value |  |  |  |  |  |
| Statistic all |  |  |  |  |  |
| y |  |  |  |  |  |
| Significant |  |  |  |  |  |
| $?$ |  |  |  |  |  |
| $?$ |  |  |  |  |  |



NAME:

Sample: $\qquad$

| $\%$ of square covered |  |
| :--- | :--- |
| White | Red |
| 1 | 1 |
| 2 | 2 |
| 3 | 3 |
| 4 | 4 |
| 5 | 5 |
| 6 | 6 |
| 7 | 7 |
| 8 | 8 |
| 9 | 9 |
| 10 | 10 |
| 11 | 11 |
| 12 | 12 |
| 13 | 13 |
| 14 | 14 |
| 15 | 15 |
| 16 | 16 |
| 17 | 17 |
| 18 | 18 |
| 19 | 19 |
| 20 | 20 |


| White | Red |
| :--- | :--- |
| 21 | 21 |
| 22 | 22 |
| 23 | 23 |
| 24 | 24 |
| 25 | 25 |
| 26 | 26 |
| 27 | 27 |
| 28 | 28 |
| 29 | 29 |
| 30 | 30 |
| 31 | 31 |
| 32 | 32 |
| 33 | 33 |
| 34 | 34 |
| 35 | 35 |
| 36 | 36 |
| 37 | 37 |
| 38 | 38 |
| 39 | 39 |
| 40 | 40 |



Sample: $\qquad$

| \% of square covered |  |
| :--- | :--- |
| White | Red |
| 1 | 1 |
| 2 | 2 |
| 3 | 3 |
| 4 | 4 |
| 5 | 5 |
| 6 | 6 |
| 7 | 7 |
| 8 | 8 |
| 9 | 9 |
| 10 | 10 |
| 11 | 11 |
| 12 | 12 |
| 13 | 13 |
| 14 | 14 |
| 15 | 15 |
| 16 | 16 |
| 17 | 17 |
| 18 | 18 |
| 19 | 19 |
| 20 | 20 |


| White | Red |
| :--- | :--- |
| 21 | 21 |
| 22 | 22 |
| 23 | 23 |
| 24 | 24 |
| 25 | 25 |
| 26 | 26 |
| 27 | 27 |
| 28 | 28 |
| 29 | 29 |
| 30 | 30 |
| 31 | 31 |
| 32 | 32 |
| 33 | 33 |
| 34 | 34 |
| 35 | 35 |
| 36 | 36 |
| 37 | 37 |
| 38 | 38 |
| 39 | 39 |
| 40 | 40 |



