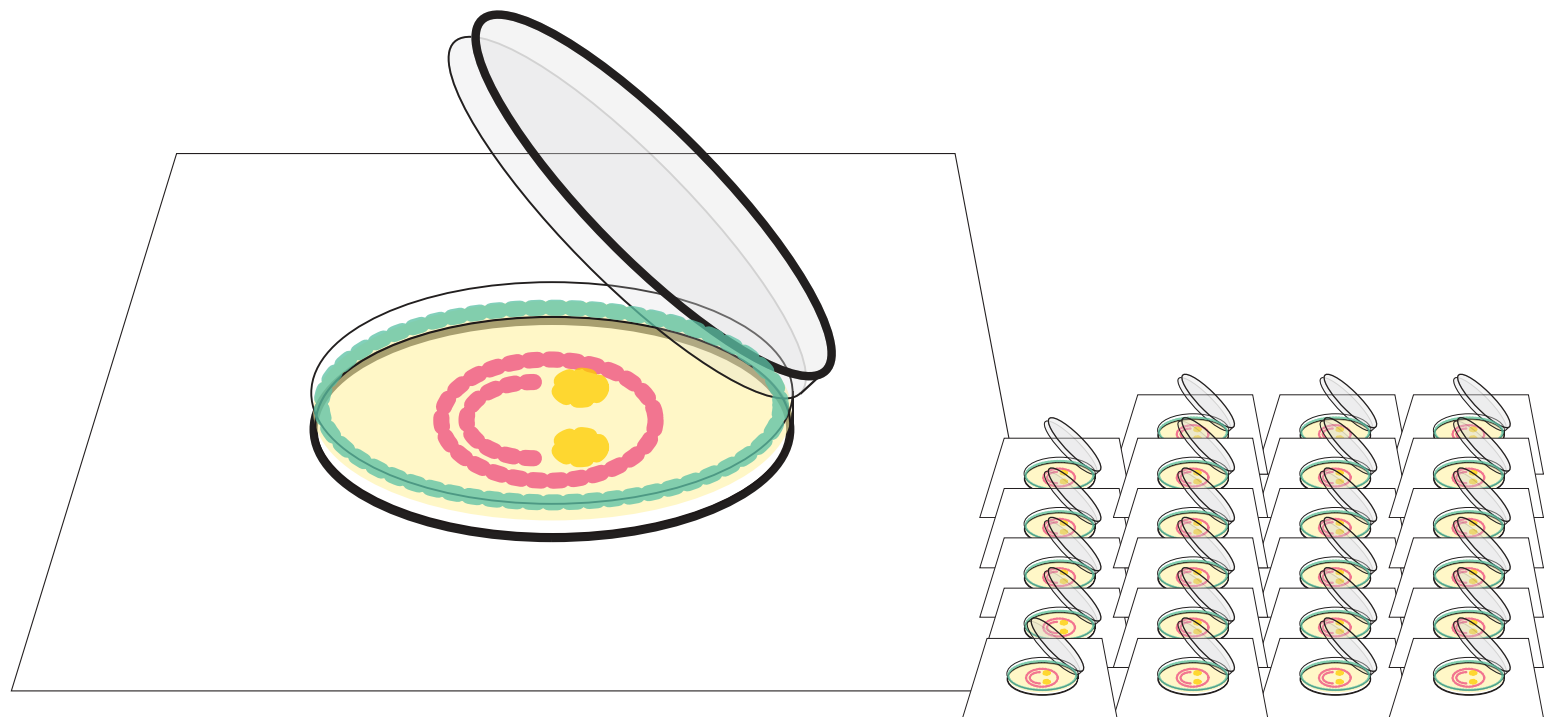


CANVAS KIT™

CLASSROOM & GROUP MANUAL

For use with the **group / large** kit size

a (2x) 45-minute class + (1x) 15-minute class periods experiment procedure



CANVAS KIT™

CLASSROOM & GROUP MANUAL

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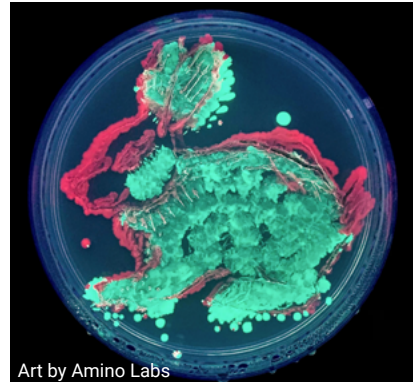
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Welcome! Let's get started



Art by Flowery Branch HS



Art by Amino Labs



Art by Flowery Branch HS

This user guide was created to help you get the most out of your Amino Labs experience. Even if you are familiar with genetic engineering, science or other Amino Labs™ products, please take the necessary time to read through this guide. This will ensure you practice safe science as well as store, use, and get the most out of your kit. It will also let you know what to do in case of a spill or accident.

In the first section, you will learn about your kit's components, how to store them before and during your experiment, as well as a few tips on activities to complete before you get your hands wet. The second section is procedural -- these are the step by step instructions on how to run your experiment. Make sure to follow our tips to ensure your best success! The third section covers "what's next"; how to keep your creations, store or dispose of any leftover ingredients and general clean up instructions. The final section is there to help you -- a glossary, troubleshooting, and our contact information.

Amino Labs is excited to welcome you to the world of the genetic engineering with the, Canvas Kit™, Engineer-it Kit™ and our entire ecosystem of easy-to-use, easy-to-succeed at products!

Following this guide will help ensure that you are getting the most out of your current and future experiences to keep on making new creations with DNA. Have fun!

Practicing safe science

Genetic engineering and life sciences are safe activities when you follow simple guidelines. Read on to ensure you adopt safe practices.

The kit in your hands contains only non-pathogenic ingredients. These are part of the biosafety Risk Group 1 (RG1) (Biosafety Level 1). This is the most benign level and therefore the safest: with these ingredients, no special containment or training is required in North America*. However, you must follow these safety guidelines for your safety and the success of your experiment(s)!

We recommend the system and kits for ages 12+, under adult supervision, and 14+ with or without supervision.

We recommend that an adult empties the discard container and that the cleaning instructions be strictly followed for safety and experiment success. Make sure to store the ingredients in accordance with the instructions found in this booklet. Eye-wear is not provided but can be worn.

- **Do not eat or drink near your experiments.** Keep your experiment at least 10 feet from food, drinks, etc. Under no circumstances should you eat any of the ingredients.
- **Immunocompromised persons:** While the ingredients in these kits are non-pathogenic, some persons, such as immunocompromised persons, can be affected by large numbers of bacteria and should talk to their doctor before doing any experiment.
- **Wash your hands before and after** manipulating your experiment, the ingredients, or the hardware.
- **Wear gloves**, even when cleaning your station or handling the consumables (petri plates, loops, etc). This will protect you from your experiment, and your experiment from you. Any latex, nitrile, or general purpose gloves you can find at the pharmacy will do. Also, after you put your gloves on, be aware of what you touch. Try not to touch your face, scratch itches with your gloved fingers!
- **If using the DNA Playground™ or BioExplorer™ place it on a stable work surface.** Keep it level at all times.
- **Clean up your station, spills and work surface before and after use.** Use a 10% solution of chlorinated bleach generously sprayed onto a paper towel and rub onto any contaminated surfaces. (Careful! This can discolor your clothes). A chlorinated spray cleaner also works.
- **Find a container to hold the inactivation bag where you will discard used consumables.** An old 1L yogurt container, large plastic cup or the like will do. Used consumables will be loops, any tube or used petri dish.

You can download a biosafety poster for your space from www.amino.bio/biosafetyinaction

If you would like to do a short online lab safety course for your edification, we recommend a Government of Canada course: www.amino.bio/biosafety

How will students learn?

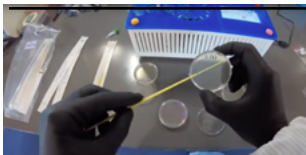
Learning and prototyping with genetic engineering and cells is becoming accessible to newcomers ages 12+ thanks to dedicated scientists and kits such as the one you are about to use!

One of the easiest ways to learn a new science, hobby or topic is by trying it hands-on. Amino Labs kits make it easy to do science by following the instructions in this booklet. Everything you need is included; each ingredient in the kit is pre-measured and labeled for a beginner-friendly experience. Our all-in-one DNA Playground minilab (mini-laboratory) decreases setup time, mess, guesswork and the need to collect and calibrate multiple machines. The included instructions should be easy-to-follow for everyone but may contain some new terms for which we have added a glossary at the end.

We also have additional resources to help students and teachers go further:



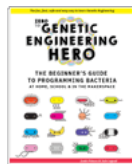
An essential addition to our ecosystem are the free **Virtual Bioengineer™ simulations** developed with the educators at the Biobuilder Educational Foundation. These simulations are 20 minutes guided experiences that make it easy to practice using a DNA Playground™ and experiment kits beforehand. The simulations includes additional information on the manipulations and a more in-depth look into the kit components. We recommend it strongly! Complete online at www.amino.bio/vbioengineer.



View **video tutorials** at youtube.com/c/AminoLabs.



Would you like for an Amino Labs team member to guide you through your journey? Try the **Cyber Workshop & Tutoring**, a 3-day+ experience completed via video conferencing. www.amino.bio/products/cyberworkshop.



Are you interested in teaching the theory behind the experiment? In going deeper on the science, learning pro-tips and eventually moving onto advanced genetic engineering? The **Zero to Genetic Engineering Hero book** is for you. Find out more at www.amino.bio/book

Discover your Canvas Kit™



The Canvas Kit™ lets you use your colored engineered bacteria to create living paintings! By following the experiment instructions on the next pages, you will create selective agar petri dish “canvases”, use the bacteria “paint-brushes” to create your living art on the agar surface and incubate over 24 to 48 hours to let your creativity grow!

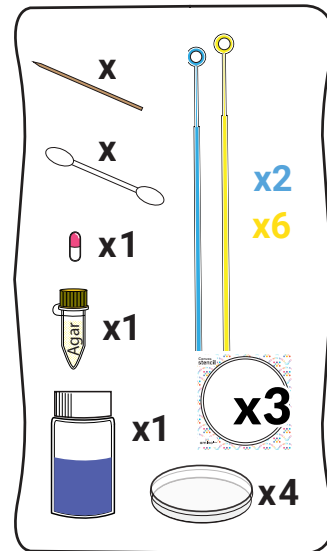
While the Canvas Kit comes in Individual Size or Group/Classroom Size, these contain the same ingredients, in different quantities. This classroom-specific manual is aimed at teachers using the group size of the Canvas kit. The group kit is perfect for a class of 24 divided into groups of 3. Diagrams on page 12 show how the kit breaks down into student packs and how to divide your students into groups.

If you are teaching or doing the exercise as a small group or alone, we have a manual available for you. Visit www.amino.bio/instructions to download the INDIVIDUAL version of the manual for the Canvas kit.

Group size - Who gets what?

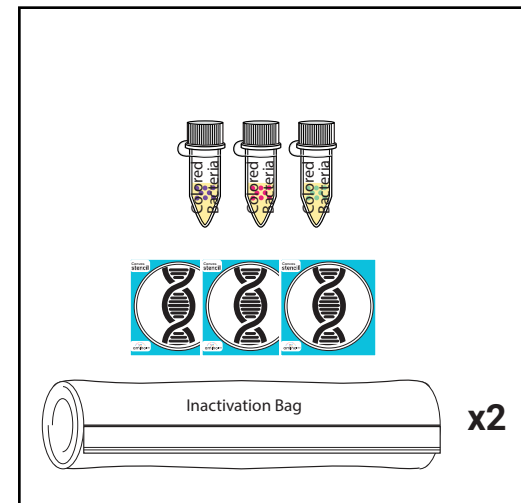
Ideal for 8 student groups with three or less per group*. The Canvas Kit™ for groups contains eight student group packs which can each make three living art pieces, and one bacteria paint painters palette. It includes a shared materials bag with the colored bacteria paint, and the inactivation bags to be shared by all groups.

Student group pack



x 8

Shared materials



* How many living paintings will be created?

Using the standard instructions 24 art pieces are possible with this kit. BUT! You can modify this to suit your needs: In total, this kit creates 32 LB agar petri dishes. As a standard, 8 of these LB agar petri dishes will be used to grow the bacteria paint to make painting palettes, resulting in 1 painting palette per student group, or 1 per student pack. 24 of the LB agar petri dishes will be used to paint the bacteria art, 3 paintings per student groups, or 3 per student pack. If you are 25 to 30 participants, you can opt to have only a few of the student groups prepare the painting palettes so that more LB agar petri dishes can be used for paintings. For example, with a group of 30 students, only 2 student groups would create painting palettes. 2 painting palettes can suffice, as long as everyone remembers that they only need to touch the bacteria paint to transfer it to their painting. No need to scoop up large amount of bacteria paint from the painting palette!

Kit components

In each student pack:



Sterile Water: Sterility is critical when genetic engineering. This sterile water bottle contains distilled water sterilized in an autoclave to ensure there are no contaminating organisms present. This 50 mL volume, when used with LB agar powder is enough to make 4 LB agar plates.¹



Small Blue Loops: Small inoculating loops are used for transferring 1 uL of liquid and other tasks. These replace costly pipettes.



Large Yellow Loops: Large inoculating loops are used for transferring 10 uL of liquid and other tasks. Yellow loops are great for spreading out bacteria after a transformation.



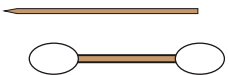
Petri Dish / Plate: 6cm Petri dishes are large enough for this lab experiment and help save on the cost of reagents and reduce waste.



Image stencil: These stencils can be used as your bioart image. Place the stencil under the petri dish to trace.



Blank stencil: These stencils can be used to draw your bioart image before tracing it on the agar. Place the stencil under the petri dish to trace.



Paintbrushes: Sterile swabs and picks to help you paint your bacteria on the agar.



Day 1 bag

Agar Powder: This LB agar powder is industry standard. Each tube of LB agar powder can make 50 mL of molten LB agar (3.5% w/v). Agar is both the surface the bacteria grow on and the food they eat to grow.¹



Antibiotics/Selection Marker: Amino Labs' proprietary antibiotic delivery system helps stabilize antibiotics for shipping and long-term storage. These capsules have a measured amount of antibiotics for 50 mL of molten LB agar. The amount of antibiotics included in the capsule is 1000 less than a standard dose for a toddler.¹

In the shared resources bag:



Inactivation Bag: A heavy duty bag to put opened tubes, used loops and petri dishes in. After the experiment is completed, simply add bleach and water to the bag to inactivate all the material and practice safe science as per *Storage, disposal & clean up* Instructions.



Bacteria paint bag

Colored bacteria paint (optional / colors vary): These colored bacteria are naturally colorful or engineered to be colorful. In either case, they are lab-strains that are non-pathogenic.

¹ For education purposes only.

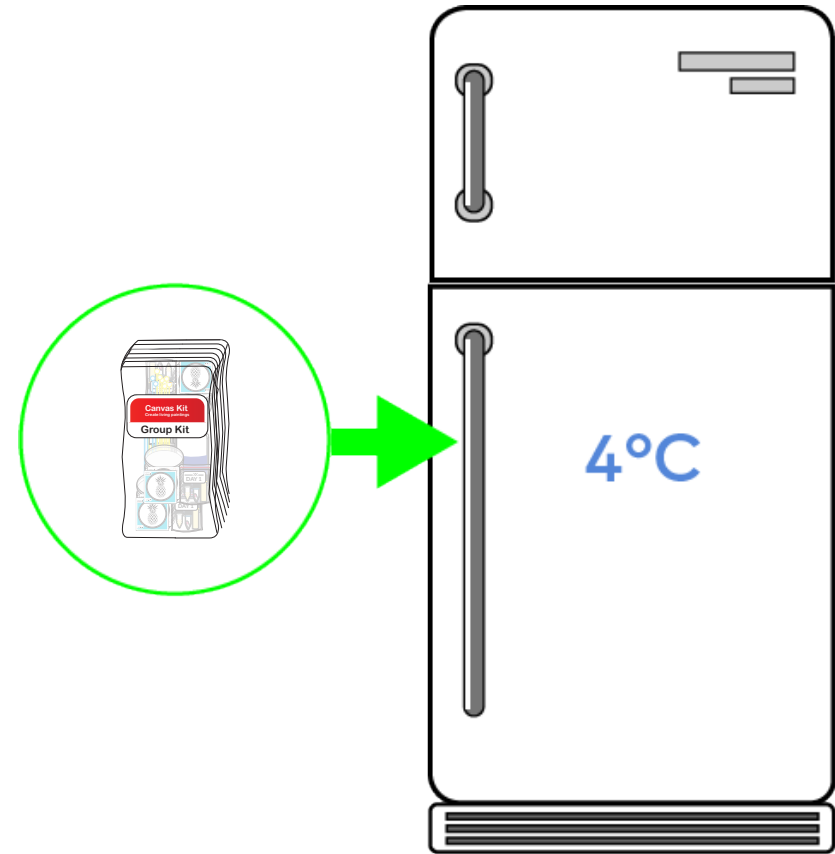
Unpacking and storing kits

For a better shelf life and successful experiments, place your Canvas Kit™ in a standard refrigerator at around 4°C.

If you can fit the whole pack, go ahead and store it all in the refrigerator. If you need to save space, the most important bag to refrigerate is the Shared Resources bag. Note that you need to keep your students packs and shared resources bags from different types of kits separate. For example, if you opened up your Canvas kit(s) and also have Engineer-it kit(s), or other group kits, you will need to identify them as the contents will be different.

If your refrigerator is not a science-only refrigerator, we recommend placing your science experiments inside a sealed plastic container before placing them in the refrigerator, especially once your kit is open.

Do Not Freeze your kit!



Technical specs

Growth plates: 6 cm petri dishes
Selection/Antibiotic: variable

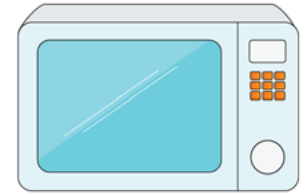
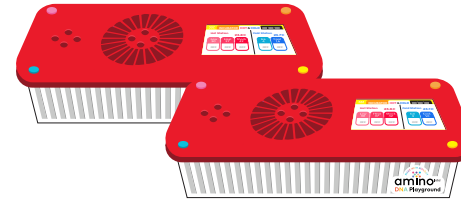
Bacteria paint: engineered *E. coli* stab

Solid growth media:
LB agar powder (1.6 g)
50 mL sterile water

Necessary equipment

For Best results:

- DNA Playground™
(One DNA Playground Classroom size per up to 3 student groups or one DNA Playground Home size per student group)
- Microwave



Alternative solution:

- **Microwave**
- **Timer**
- **Incubator** (for 37°C) : This will replace the **Incubator** set to "37". This will replace the Incubator set to "37". If you do not have an incubator (biology or egg one, as long as they set to 37°C), you can create one using an online tutorial (ex: www.instructables.com/id/Low-cost-and-accurate-incubator-for-DIY-biology/.) If you have neither incubator or DIY version, you can try incubating the cells in a resealable bag in a warm environment. Your yield won't be as good as with an incubator but should work. Note that it will take a few more days to see results and the bacteria colors will not be as bright.



Necessary safety supplies



Disposable container 500ml-1L

to hold tubes, loops and other contaminated waste (e.g., yogurt container, plastic cup). 1 per station

Latex or nitrile gloves

like the ones found at a pharmacy. 1 pairs/student if students keep & reused each day, or 3 pairs/student if not saved & reused.

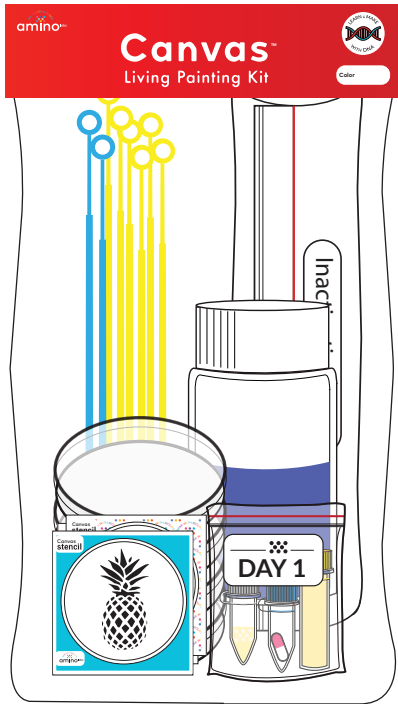
Chlorinated bleach spray

1 to share in the classroom (or you can mix a 10% solution: 1 part bleach to 9 parts water in a spray bottle)

Bleach ~500 mL

to inactivate all the experiment materials at the end of the experiment.

3-class period timeline



Canvas Kit

Day 1

Create painting palette(s)

Each student group or individual makes LB agar plates with antibiotics (selective plates) and streaks each of the colored bacteria paints from the tubes included in the kit onto one of these selective plates. This amplifies the colorful bacteria so there is enough to paint with. Incubate it so that it becomes the painting palette.

(60 minutes)

Day 2

Paint your bioart

Each student or individual can use a blank stencil to draw their picture that will become the bioart. Set the stencil under the petri dish in order to trace it using the bacteria paintbrushes and the painting palette. Incubate the petri dish(es).

(30+ minutes)

Day 3

See your bioart

View the living art grow and change color over the next 24-72 hours. Use natural and UV light to see the different colors and photographs!

(10+ minutes)

The Canvas Kit™ takes 2 days of hands-on activity to complete, and 24 to 72 hours to see results. 5 activities make up the Canvas Kit experiment:

1. Make selective LB agar plates

Day 1, 20-35 minutes

2. Streak colored cells to make enough paint

Day 1, 20 minutes, incubate 24 to 48 hrs*

*If you need to incubate your bacteria paint for longer than 48 hrs (ex: over the weekend) you can incubate it at 30°C instead of 37°C. This is only okay in the Canvas kit!

3. Stencil art on paper

Day 2, variable time

4. Paint with Bacteria

Day 2, 20+ minutes, 24-72 hrs incubation

5. View results

Day 3, 20 minutes

Recommended pre-labs

Amino Labs has some resources that should be used by your students before they complete the hands-on experiment to maximize their understanding and success. These pre-labs are meant to ensure your students know, understand, and complete all the experiment steps. Completing the pre-labs also minimizes the number of questions your students will have during the hands-on experiment.

1. Virtual Bioengineer Simulator - Canvas Kit Edition

www.amino.bio/pages/vbioengineer_canvas

This free simulator walks your students through the entire Engineer-it Kit's materials and procedure. The students can complete the simulator as homework or in class with the use of the school's computer lab or the student's laptop computers. The simulator takes approximately 25 minutes to complete. It is also common to project the simulator and complete it as a group during class or as a review if the students have completed it as a homework assignment.

2. Canvas Kit experiment procedure - a quick test

www.amino.bio/canvas-pre-test

A short activity to test whether your students read the experiment procedures and are ready to start the experiment. The students can complete the test online on a computer or mobile device or you can print it out.

2 key pitfalls to avoid!

In the next pages are detailed, step-by-step instructions to complete the experiment. These include instructions to prepare the classroom and the students' instructions. **Please make sure the students read all the steps before starting the hands-on manipulation;** some steps will be done in rapid sequence. The best way to ensure students success is by having students complete the recommended pre-labs on the previous page.

While all the steps outlined in the experiment protocol are important and should be followed as described, the **MOST IMPORTANT** considerations for success are:

1. In Step 1: When making the LB agar, make sure that the water is boiling before adding the agar powder. **Students have to see the water bubbling!** Caution, the bottles will be hot!

2. In Step 3: **Before painting the art or flipping the painted petri dish upside down for incubation, make sure the agar is dry and there is no condensation.** Otherwise, the bacteria paint may touch the condensation/water on the surface of the agar and spread around the agar. This means the painting will turn out very blurry!

Teacher Experiment Setup

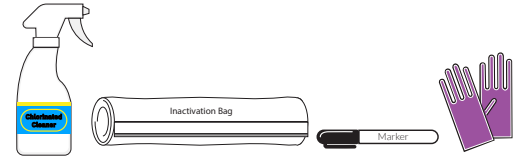
0. Prepare your classroom space

Goal Set yourself up for success.

Materials from your kit
(1 per group) Student packs
Shared materials bag

Materials not in your kit
(1 per table) Discard container
Chlorinated bleach spray or wipes
(1 per table) Permanent marker

(1) Paper towels
(1 pair per student) Gloves



Make sure the class has access to a microwave before starting!

0.0 Have students download/print the manual and read *Practicing safe science*, the *2 pitfalls to avoid*, the *Student experiment protocol* (including the *Checkpoints*), and the *Glossary* pages.

0.1 Set down the DNA Playground(s), or other lab equipment (37°C incubator) on or near the students work stations. Make sure the equipment is level and on a stable surface. Refer to the instruction manual to make sure you know how to use your equipment safely. For this kit, you can also incubate at room temperature - no equipment needed. Note that room temperature incubation will add more incubation time to your experiment.

0.2 Set one discard container per work station (as per the *Necessary safety supplies* page).

0.3 Set one Student Pack and one permanent marker (sharpie) per student-group area. Keep the shared materials in a common area so all students can access them on step 2.

0.4 Ask the student to use the discard container to dispose of:

- any used inactivation loops, paintbrushes
- empty tubes (agar, antibiotics tubes, etc...)
- any gloves that have touched bacteria.

Paper, plastic packaging and gloves that have not come into contact with bacteria should be disposed in the regular garbage or recycling bin. After each day's experiment or at the end of the entire experiment, have students pour the content of their discard container into an inactivation bag. Follow the instructions at the end of the experiment to inactivate the contents.

0.5 Ask the students to put on their gloves.

0.6 Have the students wipe down their work surface with chlorinated bleach spray, wipes or 10% bleach solution. Do not have them spray bleach solution directly on the DNA Playground.

0.7 After the students complete the experiment, follow the *Storage, discard & clean up* procedures with them.

If you are saving the tubes of cells for a future experiment, place back in their ziplock bag after use and refrigerate. We recommend you use a sealed plastic container to store all your experiment materials inside a refrigerator if you also use this refrigerator to store food or drinks. If you are not saving them, place the open tubes in a discard container and dispose of them after all the student-groups have used them.

Student Experiment Protocol

1. Creating selective LB Agar Plates Day 1, 25 minutes

Goal Create selective LB agar plates.

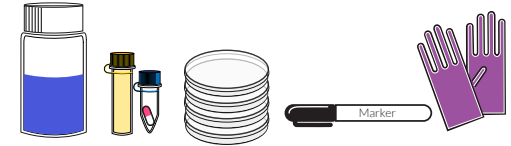
Materials from your kit

(1) 50 mL sterile water
(1) LB agar powder

(1) antibiotic pill
(4) 6 cm petri dishes

Materials not in your kit

(1) Sharpie marker



Prepare

1.1 Use a sharpie-type pen to label the bottom of the petri dishes (the side with the star-shaped ring) as follows:
4x S. (for selective) + Add [your initials] if doing this in groups with multiple kits.



Mix the Agar

1.2 Unscrew the lid from the sterile water bottle and keep it loosely on top of the bottle to prevent any contaminants from entering the water, but allowing air to escape. This will prevent pressure build-ups.

1.3 Place the bottle in the microwave and heat the water **until you see it boil**. You can use 45 seconds as your starting time but you have to see a rolling boil where many bubbles are rising constantly before you continue to the next step. Careful, the bottle will be hot! **!! If the water does not boil, the agar powder will not dissolve and your plates will not solidify !!**



1.4 Add the tube of agar powder to the boiling water. Careful, the water is hot! Some agar powder may “clump” around the lip of the tube due to the water evaporation. This is okay, we have accounted for this possible loss.

1.5 Microwave the water and agar powder in 4 seconds intervals until you see it boil again. Instead of a rolling boil, you will see more of a foam forming above the molten LB agar liquid. **Careful, the liquid will boil over if you microwave in more than 4 sec. increments.** After you see the liquid foaming, swirl to mix for 10 seconds.

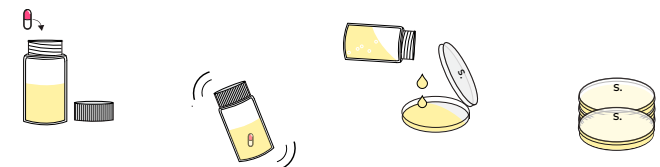


Note: If you've done the Engineer-it Kit before, note that you will not be making a non-selective plate. All 4 plates will be selective agar.

Make selective (S.) plates

1.6 Add the antibiotic pill to the bottle of agar and gently swirl for a few minutes until the contents of the pill have dissolved. Do not introduce bubbles into the LB agar: don't swirl too vigorously. The gelatin capsule may not fully dissolve. The important thing is that the contents of the capsule do dissolve.

1.7 Once the antibiotic pill is dissolved, pour the molten LB agar into the 4 petri dishes. Place the lids 3/4 of the way back on so that the agar can cool and dry (solidify).



Pro-tip: If there are water droplets on the surface of the LB agar, this can disrupt your art. Bacteria that you will

be painting with can enter a droplet and spread throughout the droplet therefore 'smudging' your art. To avoid this make sure the lid is partially over top to allow for evaporation and a dry LB agar surface.

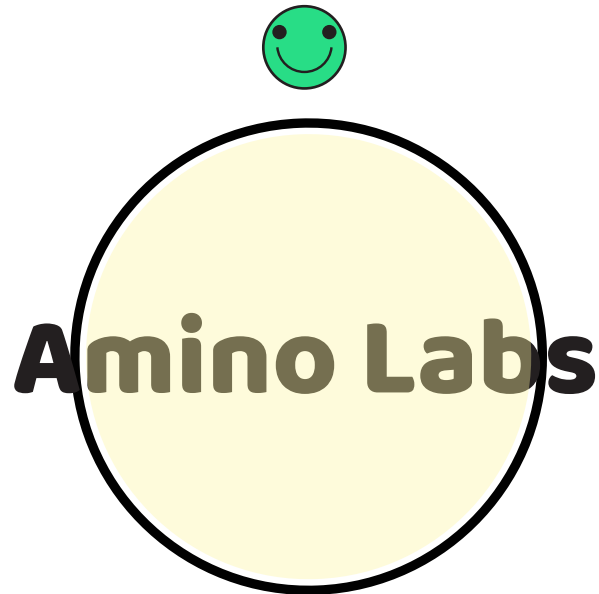
1.8 Let the LB agar harden. This can take up to 20 minutes depending on how warm and humid your environment is. You will use 1 plate in the next step. You can store the remaining 3 plates in the zip-lock bag in the refrigerator for day 2.

Troubleshooting tip

If your plates do not solidify after 30 minutes it is very likely that the water was not boiled enough to dissolve the agar powder. As a 'hack', you can pour all of the petri dish content back into the water bottle and microwave until you see it boil. Swirl to mix and re-pour your plates.

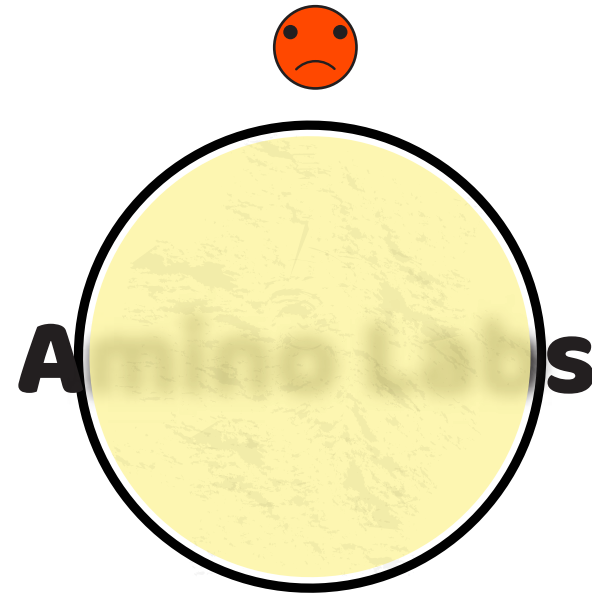
Checkpoint - Agar Plates

Use this guide to check if you are ready to move onto the next step.



A perfect Agar plate is completely clear and solid - if you set it 4" above some image or text, you should be able to read it / see it clearly.

Move on to the next step!



An agar plate that is cloudy and/or bumpy and/or soft is not ideal - if you set your plate 4" above some text or image and cannot see clearly through it, it means you needed more boiling or mixing.

Troubleshooting tip

If your plates do not solidify after 30 minutes it is very likely that the water was not boiled enough to dissolve the agar powder. As a 'hack', you can pour all of the petri dish content back into the water bottle and microwave until you see it boil. Swirl to mix and re-pour your plates.

Unfortunately, if the agar does not solidify, this means you need to halt your experiment and complete the troubleshooting guide and follow the instructions at www.amino.bio/troubleshoot

2. Create your painting palette Day 1, 25-45 minutes + 24 hours wait time

Goal Create bacteria paint painting palettes.

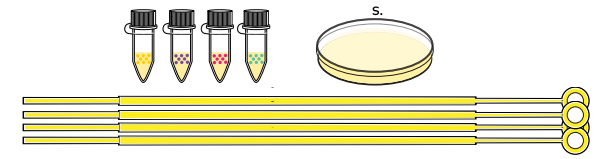
Materials from your kit

(1) Selective agar plates

(4) Yellow loops

Shared resources

Tubes of bacteria paint



Prepare

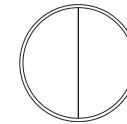
2.1 If you have an incubator, turn it on to 37°C.



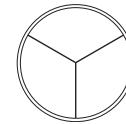
Streak

2.2 Take your tubes of colored bacteria and look at the colors you have available. Using a sharpie, divide up the bottom of your petri dish into as many sections as you have colors like you would divide a pie. For example, divide the petri dish into 2 sections if you have 2 colors, 3 sections if you have 3 colors, and so on.

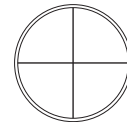
2 colors



3 colors



4 colors



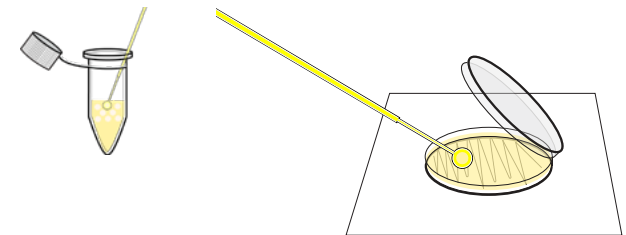
2.3 Use the sharpie to write the name or the abbreviation of the different colors of bacteria in each section on the bottom of the petri dish. One color per section. The order does not matter, as long as each color has a section.

2.4 If you have printed instructions, place your petri dish on top of one of the stencils on the next page. Choose the stencil that is divided into the same number of sections as your petri dish. If you are using online instructions, you can manually copy the right stencil pattern on a piece of paper and place your petri dish on top. For this streaking, it is not important to be precise, it is simply necessary to cover most of the surface with the bacteria.

2.5 Open one of the yellow loop by holding the straight end of it, not the loop end. Remove from the packaging. Don't let the loop end touch anything yet!

2.6 Open one of your colored bacteria tube. Dip the loop end of the yellow loop into the stab of colored bacteria.

2.7 On your petri dish, find the section that you marked for this color and, using the end of the loop you dipped in the colored bacteria, trace the zigzag line like on the stencil on the next page.



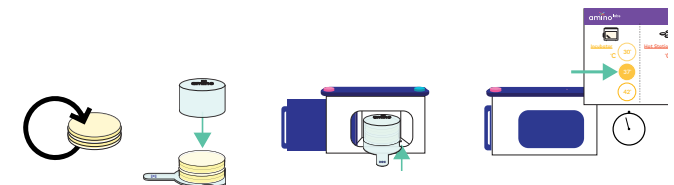
2.8 Discard the loop in your inactivation bag or discard container.

2.9 Using a new yellow loop each time, repeat steps 2.5 - 2.8 for each colored bacteria tube you have.

2.10 Close your tube(s) of bacteria. Return them to the group sharing area.

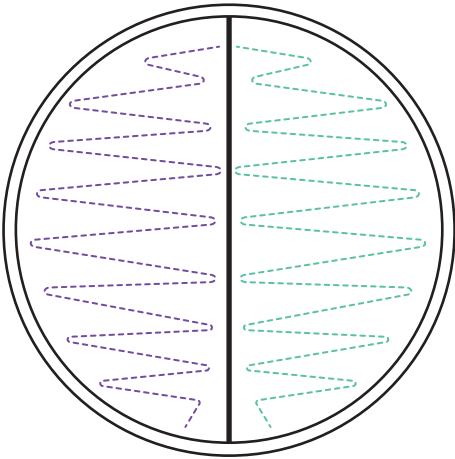
Incubate Overnight

2.11 Incubate your streaked plate **upside down** at ~37°C for 24 to 48 hours. This will be your biopaint to create your living art. (It may take longer if you do not have an incubator, up to 5 days. Note that the colors will be more pastel if you are incubating at room temperature). *If you need to incubate your bacteria paint for longer than 48 hrs in an incubator (ex: over the weekend) you can incubate it at 30°C instead of 37°C. This is only okay in the Canvas kit! * **Note: Remember to flip your plate upside down! If you have Amino Labs' minilab, remember to put the humidity chamber on top of your plate and to close and lock the incubator door.**

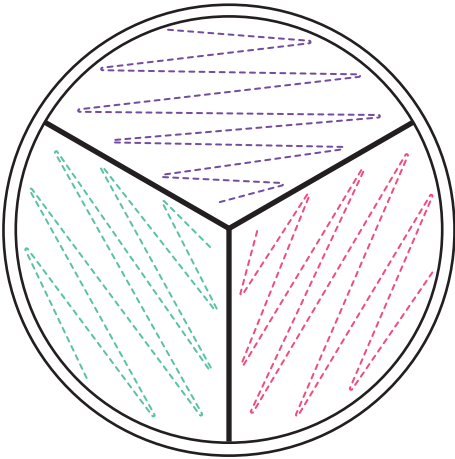


Stencils

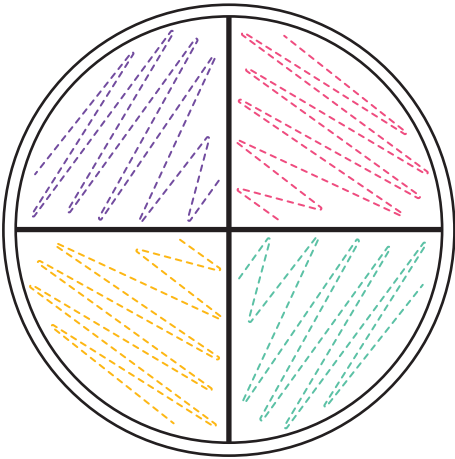
2 colors



3 colors

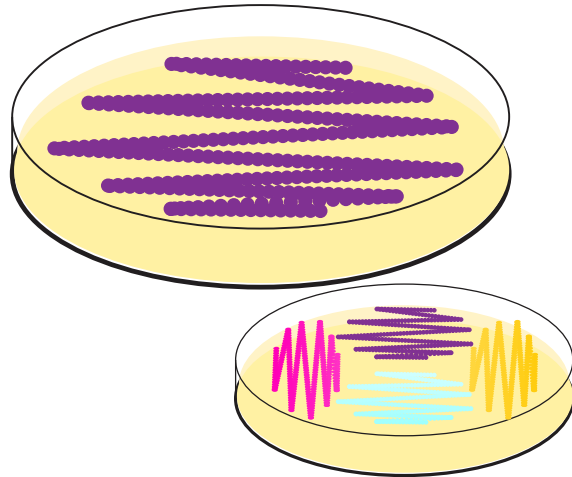


4 colors

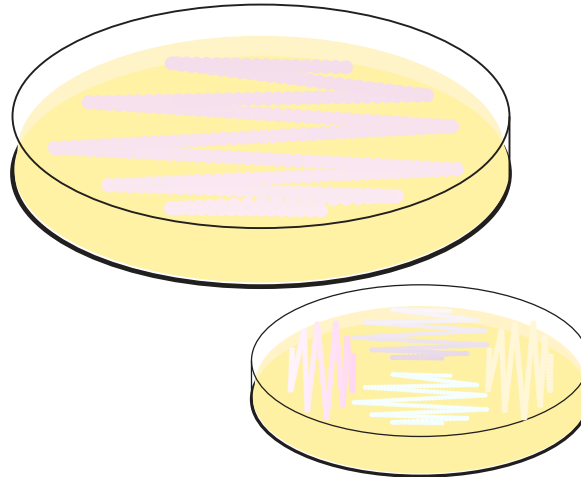


Checkpoint - Bacteria Paint

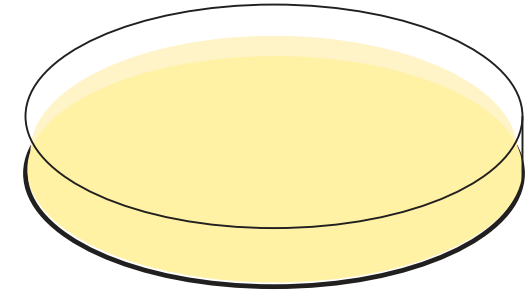
Use this guide to check if you are ready to move onto the next step.



A perfect bacteria plate has lots of brightly colored bacteria after incubation. Proceed to the next page. This suggests that you have made your LB agar petri dishes properly and have the right amount of antibiotics. **If you are incubating at room temperature, your bacteria will not become as bright as below. That's ok!**



A bacteria plate showing lightly colored bacteria after incubation requires more time to grow. Continue incubating, checking every 12hrs, until the colors are bright. If your bacteria are not changing color you may have forgotten to add the antibiotics or you re-microwave your agar which degraded the antibiotics. Contact help@amino.bio



If you see no growth on your plate:

1. If your incubator was not at 37°C or is homemade, incubate for another 24hrs.
2. If you are certain you incubated at 37°C, or incubated for 48hrs and still have no colonies, you might not have had cells on your loop when you streaked. Repeat Step 2: Grow your bio paint on the plate.
3. If you still have no colonies after repeating Step 2, contact us at help@amino.bio, and we will help you succeed.

What can you expect the colors to look like?

Under regular light

Cyan



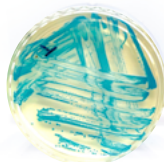
Purple



Magenta

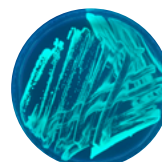


Teal

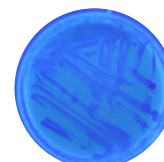


Under UV/blacklight

Cyan



Purple



Magenta



Teal



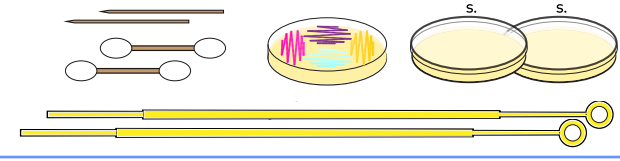
3. Paint with bacteria! Day 2, 30-60 minutes + 24+ hours wait time

Goal Create living paintings

Materials from your kit

(1 per student) Selective Agar petri dish Biopaint plate from yesterday
Loops

"Paintbrushes"

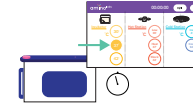


Prepare

3.1 Make sure you have colored bacteria on your biopaint petri dish from the prior day. If colors have not appeared yet, wait longer, up to 48 hours. Once you have colors, take your petri dish of biopaint from the incubator. (You can also use your freshly engineered bacteria if you completed an Engineer-it Kit previously).

3.2 If you have an incubator, turn it on to 37°C.

37°



Paint!

3.3 Using the blank stencils in your kit, sketch your art piece for each petri dish your are painting. You can also use the canvas stencil that already has an image included in the kit.

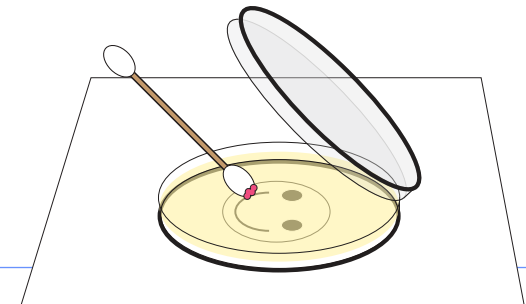
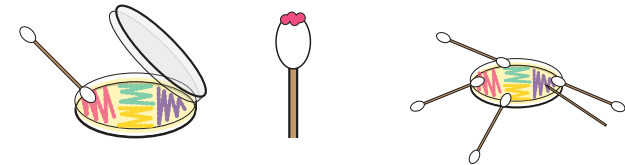
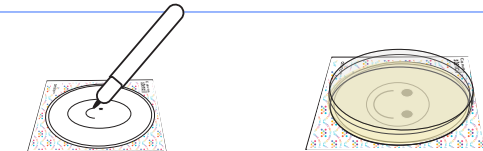
3.4 Set one of your selective petri dish canvas on top of your sketched stencil or the image stencil from the kit.

3.5 Using the yellow loops, the blue loops and the bacteria "paintbrushes", paint your art onto the agar by dipping into the colored bacteria from the biopaint petri dish and tracing your image, gliding on top of the agar. The agar is like a Jell-O, be careful not to puncture it as you paint.

Notes: Assign a bacteria color to each q-tip and/or toothpick as you only have a few of these. Set them down on the edge of the bacteria painting palette when you are not using them until you have completed your art.

You will not see the bacteria appear right away, but you may be able to see a "wet" trace where you have painted on top of the agar. **You only need to dip into the colored bacteria on the painting palette once to collect bacteria paint.**

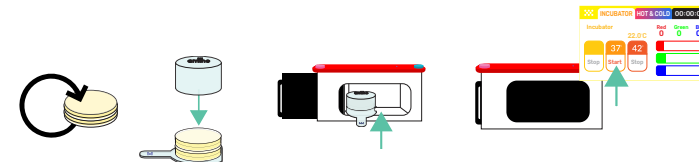
3.6 Return any unused canvas and the painting palette petri dish to the ziplock bag and refrigerate.



Incubate

3.7 Incubate your art canvases **upside down** at ~37°C for 24 to 72 hours (it may take longer if you do not have an incubator). *If you need to incubate your bacteria paint for longer than 48 hrs in an incubator (ex: over the weekend) you can incubate it at 30°C instead of 37°C. This is only okay in the Canvas kit! *

Note: You must flip your plate upside down so that the agar is up and the lid down. If you have Amino Labs' minilab, remember to put the humidity chamber on top of your plate and to close and lock the incubator door.

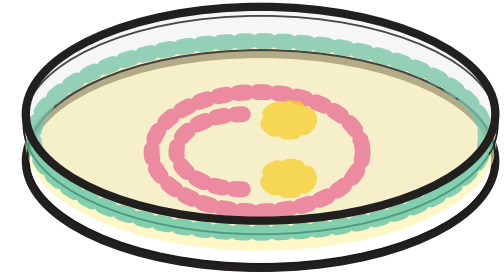


4. Did your living art grow? Day3+

Goal Verify if your bacteria paintings grew

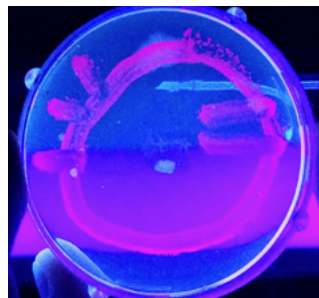
You should see your living painting appear over the next 24 to 72 hours! Keep an eye out, and your camera ready to document. **Congratulations!**

If there are any unused petri dishes left you can repeat steps 3.1 - 3.8 for those canvases or keep them in a bag in the refrigerator for up to a month. If you see any unexpected growth on these, follow the inactivation instructions.

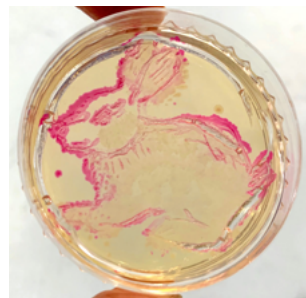


Note:

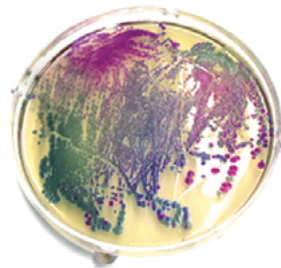
If you cannot see any growing cells at all after 48 hours of incubation at 37C, your experiment may have failed. See our troubleshooting guide at the end of the manual, compare results with your group, if applicable, or [contact us](#) with photos of your result and any documentation of your process so that we can help you succeed in the future. Make sure, if possible, to also review the video tutorials on the youtube channel ([youtube.com/c/aminolabs](https://www.youtube.com/c/aminolabs)) to see if you missed any steps!



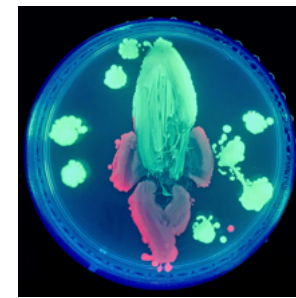
Abstract
Jim Blake



Bunny
by Amino Labs
@aminobiolab



Abstract
Neuva School
@nuevaschool



Space rocket
Teacher at an Amino Labs event
@aminobiolab

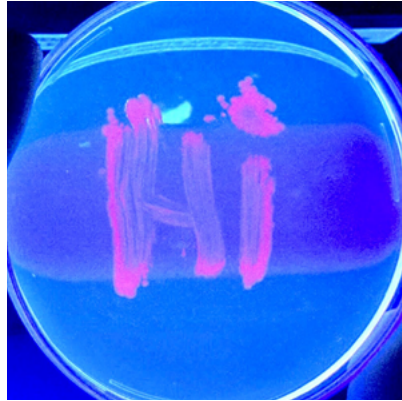


Darwin
Dr. T. Ryan Gregory
Guelph University

CONGRATULATIONS



Smiley, Anonymus



Hi, by B.L.



Portrait by Amino Labs

You have now joined the global community of bioartists! Happy with your artwork? There are many opportunities to share it online, exhibit it in your community and even participates in contests and artist communities on the web!

Share your results with friends and our growing community. Find us on Instagram, Twitter, and Facebook @aminobiolab

Don't forget, you can preserve your bioart with our *Keep-it Kit™*. For now, let's make sure you dispose of and store your remaining material correctly.

Storage, Disposal, Clean Up

After everyone sees their results, all experiment Petri dishes, tubes of cells, loops should be in the discard containers. Disposing of experiment materials is an integral part of the experiment. **Always wear gloves for cleanup!**

A. Preserving Petri dishes: If you want to preserve the living paintings or experiment results in Petri dishes instead of disposing of them, use one of our Keep-it Kits. This will help you maintain the petri dish by pouring a special resin on top. If you do not have Keep-it Kits on hand but will be getting one soon, keep the Petri dishes you want to preserve in a zip-lock bag in a cool area and out of sunlight in the meantime. You can refrigerate it to keep it “fresh” for up to a month.

B. Reusable materials: If you have DNA in your kit, it can last up to 6 months when stored in a refrigerator. If you wish to keep it, store it in a zip-lock bag inside a sealed plastic container in a refrigerator away from food items. If you do not wish to keep it, add to an inactivation bag. Make sure the lids are separate from the tubes so that the inactivating liquid can get inside. If you see any mold or unknown bacteria growing on any material at any point, immediately inactivate them by using a solution of bleach water. Follow the inactivation instructions below. If you are out of inactivation bags, use a sturdy ziplock type bag or disposable container with a lid. Always wear gloves when handling experiment materials and cleaners!

C. Unused ingredients: If you did not use all the agar Petri dishes you poured, store these for later use. Store them in their ziplock bag within a sealed container in the refrigerator for up to a few months. Keep them away from food items. If you see any mold or unknown bacteria growing inside, then you should always immediately inactivate the Petri dishes.

D. Inactivation: Dispose of bacteria, agar, tubes, loops, paintbrushes, Petri dishes, contaminated gloves, and other non-paper material from the discard containers by having the students transfer it to an inactivation bag. Remind students that any paper packaging like loop wrappers, plastic bags, and gloves that have not touched bacteria go in the regular garbage or recycling.

Make sure all the tubes have their lids off once in the inactivation bag and add a solution of 1 part bleach to 4 to 6 parts water to the inactivation bag. Close the bag and let sit for 24 to 48 hours before discarding the liquid in the toilet and the solids & bags in the garbage. Step-by-step instructions are on the inactivation bag and in an Inactivation video on youtube; youtube.com/c/AminoLabs.

Spray some chlorinated bleach cleaner in the discard container(s) once emptied. Let it sit for an hour before wiping down. You can wait to wipe it down until you empty out your inactivation bags the next day.

E. Clean your workspace: Use a chlorinated spray cleaner, wipes, or a solution of 1 part chlorinated bleach to 9 parts water to wipe down your work area and equipment. You can wipe down the minilabs with this solution and follow it with an eyeglass or window cleaner to remove the inevitable streaking from the bleach cleaner. Never use rubbing alcohol (isopropyl alcohol) on the DNA Playgrounds.

Glossary

Agar: is a Jello-like substance that serves as a growth media for bacteria. It is mixed with our bacteria's favorite food: Lysogeny broth (LB). LB is made up of yeast, vitamins, and minerals. LB can also be found liquid-form.

Antibiotics: When you transform bacteria, they will become resistant to a type of antibiotics no longer used in hospitals. This antibiotic will be mixed in with the agar and LB so that, as you incubate your culture, only transformed bacteria will grow. This is called a "selection marker".

Buffers: Buffers are saline solutions that help, in this case, open up the cell membranes so that they may take up new DNA.

Cells: Cells are tiny, living units that function like mini-factories. Bacteria are single-celled organisms (unicellular) microorganisms. They are different from plant and animal cells because they don't have a distinct, membrane-enclosed nucleus containing genetic material. Instead, their DNA floats in a tangle inside the cell. Individual bacteria can only be seen with a microscope, but they reproduce so rapidly that they often form colonies that we can see. Bacteria reproduce when one cell splits into two

cells through a process called binary fission. Fission occurs rapidly, in as little as 20 minutes.

Competent Cells: Since DNA is a very hydrophilic molecule, it won't normally pass through a bacterial cell's membrane. In order to make bacteria take in the DNA plasmid, the cells must first be made "competent" to take up DNA. This is done by creating small holes in the bacterial cells by suspending them in a solution with a high concentration of calcium (the transformation buffer). DNA can then be forced into the cells by incubating the cells and the DNA together on ice, placing them briefly at 42°C (heat shock), and then putting them back on ice. This causes the bacteria to take in the DNA and is called "Transformation".

DNA: The DNA is the set of instructions that tell the cell how to function like a computer program tells your computer what to do.

DNA plasmid: A plasmid is a small circular piece of DNA (about 2,000 to 10,000 base pairs) that contains essential genetic information for the growth of bacteria. Bacteria share vital information by passing it among themselves in the form of genes in plasmids. By inserting a new plasmid in our bacteria, we

can get them to produce things for us, like mini-factories. In this case, we have a plasmid that encodes for the creation of colorful pigments.

Heatshock: is when the cells are moved from ice-cold to warm temperature, typically 42°C, to take in DNA plasmids more efficiently.

Inoculation: is when you introduce bacteria into a medium suitable for its growth.

Inoculating Loops: are used to transfer liquids, cells, and DNA from one vial to the next instead of traditional lab pipettes, making your job easier, and less costly. They come in different pre-calibrated sizes, so you do not need to worry about minuscule liquid volumes. They are also used to spread bacteria on an agar surface without puncturing the soft agar.

Non-Selective: A non-selective plate means that any cells/bacteria put on this agar will grow as long as they are oxygen-loving organisms (called aerobic

bacteria).

Plates (or Petri dish): A petri dish is a small plastic container used to culture (grow) bacteria in a controlled environment.

Recovery period: is the period after the heat shock in which the cells develop their antibiotics resistance and start dividing.

Selective: A selective plate means it contains antibiotics. When you insert a new DNA program into cells to make them create pigments, or anything else, you also put a “selective marker” (antibiotics resistance) inside the code. This means that only the cells that have taken up the new program will be able to grow on a plate that has the antibiotics mixed in. You only get the cells you transformed!

Transformation: See competent cells.

Troubleshooting

Here are some possible common issues:

Your agar is too wet/ doesn't solidify:

When done correctly, the agar will be the consistency of Jell-O. If it is not:

1. You likely did not heat (boil) the water before, or after adding the LB agar powder
2. You might not have added all the powder from the tube, resulting in too much water vs. LB agar powder.
3. You may not have fully dissolved the powder, meaning it cannot turn into a gel and will look cloudy. You can practice by making Jell-O! Next time heat and swirl longer to ensure the powder is fully dissolved.

You don't have any colonies and its been 24+ hours:

Don't worry, every scientist has experienced this, and it can take some practice before success.

1. Double check that your incubator is on at 37°C. If it is not, or if you are growing at room temperature, then it can take much longer to see the bacteria colonies. Keep waiting!

If you kept the second half of your recovered cells, you can pour them on your plate after 48 hours of seeing no engineered colonies grow and keep incubating.

2. You may need to try again to hone your skills. See our YouTube videos for tips and tricks on how to improve your chances of success.

Your colonies of bacteria grew, but they are the wrong color or there is mold on your petri dish:

Danger! If at the end of, or during, the incubation period your resulting bacteria/plate is: a)not the right color; b)is black when it shouldn't be, this is a sign that your culture is NOT YOUR ENGINEERED BACTERIA. You should immediately inactivate it and clean your space and unit.

To inactivate it, either add it to the inactivation bag or pour 100% chlorinated bleach into the dish, put the lid on and let it sit for 24 hours before throwing it out: The strong oxidizing environment degrades any living organisms. After 24 hours, if there are still organisms present add more concentrated bleach until it is almost full, and let stand for a further 24 hours.

There may be mold in your environment. We recommend, getting a small air purifier with a HEPA filter for the room.

Always be aware that concentrated bleach is a strong oxidizing agent and if poured on the skin can cause irritation, and on clothes remove color. Follow the safety and handling protocol on the manufacturer's label.

Find an interactive troubleshooter online at

amino.bio/troubleshoot. We recommend using it for tips, tricks and to claim your Success Guarantee Kit if you need of one.

If anything else causes you issues, please contact us : help@amino.bio

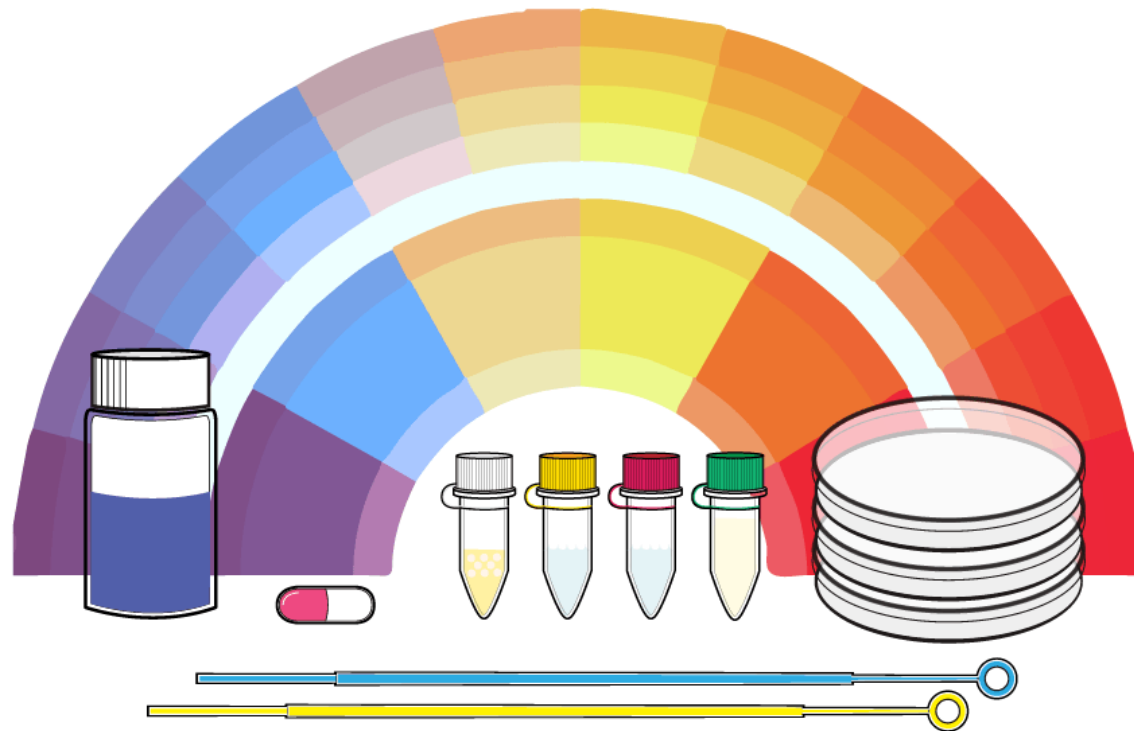


Made in Canada

All Amino Labs products, from the hardware to DNA, are invented, designed, manufactured in and shipped from our laboratory-workshop in Canada. We'd love to hear your feedback and suggestions so that we can continue to make our products better and more fitting to your needs. Answers to your questions and help are also just an email away.



Help and General inquiries: help@amino.bio
 Feedback, Suggestions, Comments: info@amino.bio



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