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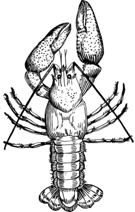
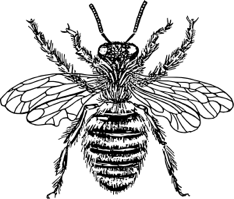
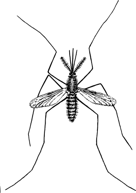
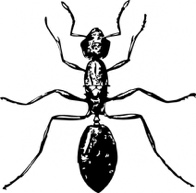
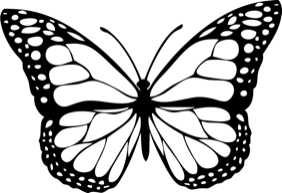
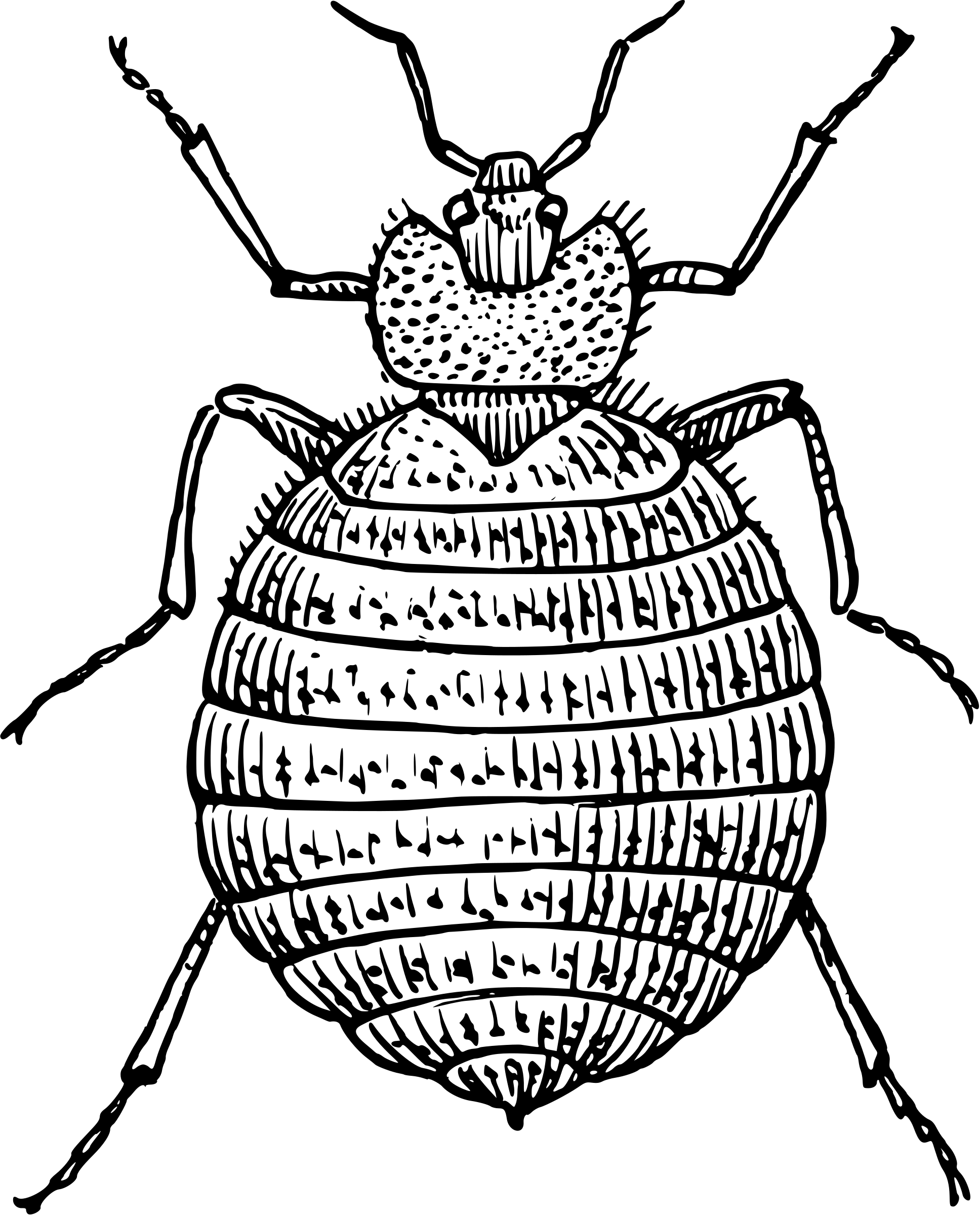
Updated: March 2024

**Lab 2: DNA Extraction**

*QIAGEN DNeasy Blood & Tissue Kit*

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**Project Guide**



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**A close up of a sign

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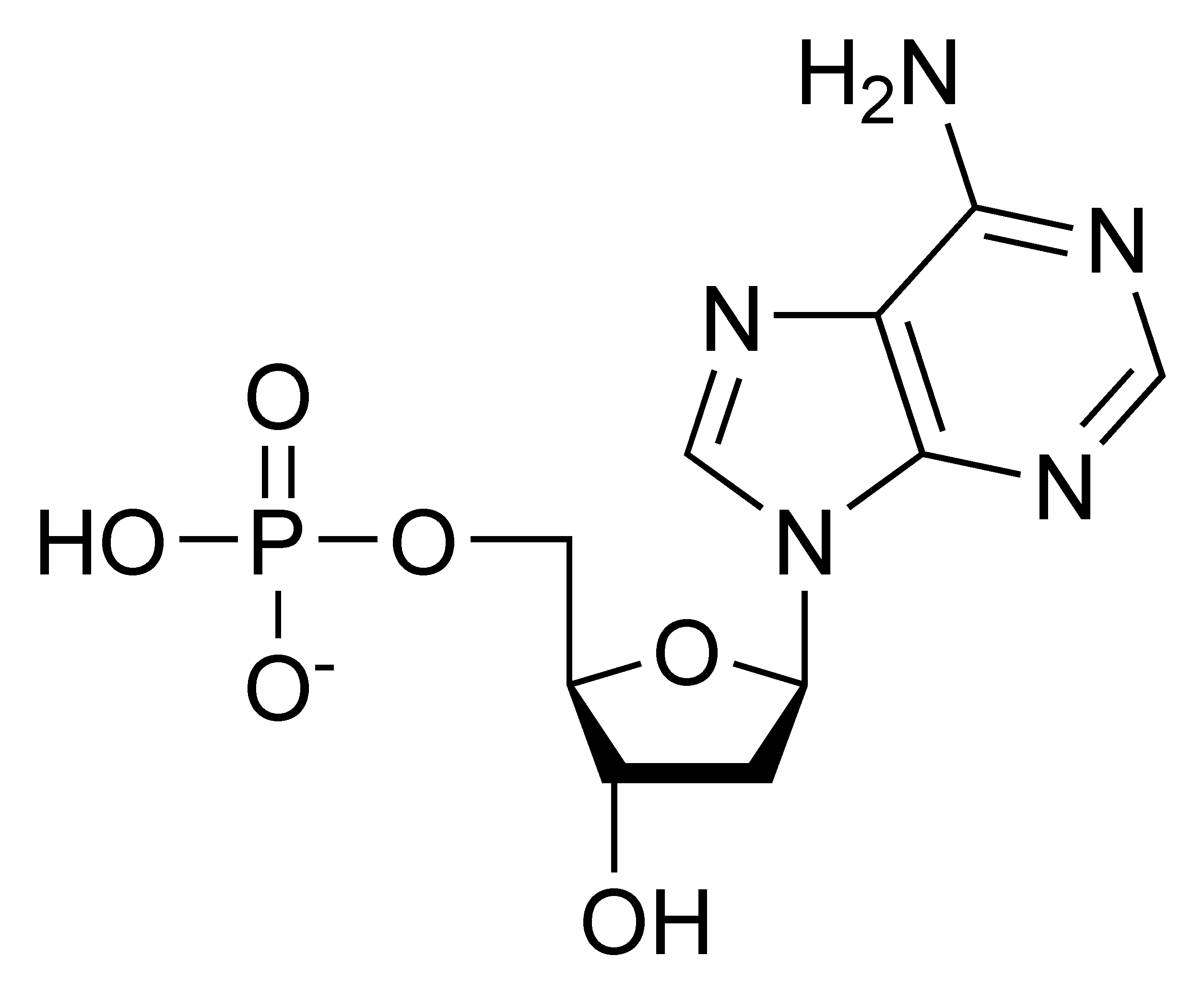
The *Wolbachia* Project: Discover the Microbes Within! was developed by a collaboration of scientists, educators, and outreach specialists. It is directed by the Bordenstein Lab.

https://www.wolbachiaproject.org

**Introduction**

**DNA Composition**

Deoxyribonucleic acid, or DNA, is a self-replicating molecule that encodes the genetic information for nearly all living organisms. DNA is comprised of monomers, called nucleotides. Each nucleotide consists of (i) a five-carbon sugar (deoxyribose), (ii) a phosphate group, and (iii) a nitrogenous base (Figure 2.1). Nucleotides are differentiated by their attached nitrogenous base: adenine (A), thymine (T), guanine (G) or cytosine (C). Adenine and guanine are classified as purines; cytosine and thymine are classified as pyrimidines. The nucleotides are linked together via phosphodiester bonds to form long strands of DNA, where the order of nucleotides along the sugar-phosphate backbone determines the DNA sequence.



**Phosphate group**

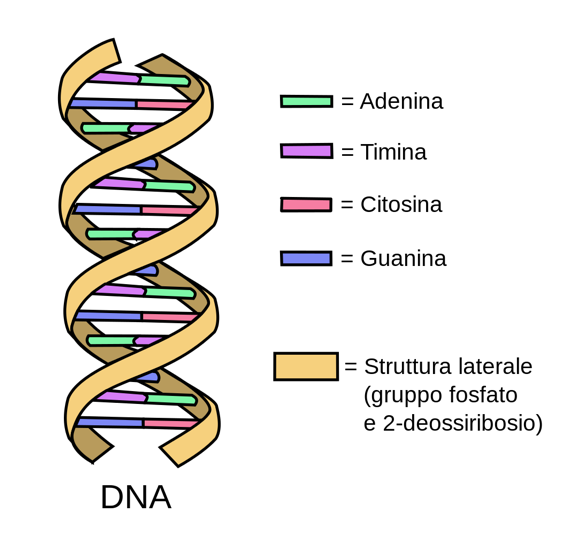
**5-carbon Sugar (Deoxyribose)**

**Nitrogenous base**

**(Adenine)**

*Image from Wikimedia Commons; available in public domain.*

*Image modified from Wikimedia Commons/Forluvoft; available in public domain.*



Adenine (A)

Thymine (T)

Cytosine (C)

Guanine (G)

Sugar- phosphate backbone

*Left: Nucleotide structure consisting of phosphate group, sugar (deoxyribose), and nitrogenous base.*

*Right: DNA double helix structure consisting of a hydrophilic sugar-phosphate backbone and hydrophobic nitrogenous base pairs.*

**Figure 2.1**

**DNA Structure**

A single strand of DNA is asymmetrical and therefore has a specific method of combining with a second strand of DNA. The sugar and phosphate group are both hydrophilic and will readily come into contact with water molecules within a cell, whereas the nitrogenous bases are hydrophobic and will aggregate together. Two strands of DNA can satisfy their hydrophilic and hydrophobic constraints by orienting their nitrogenous bases to the inside and their hydrophilic sugars and phosphate groups to the outside. The nitrogenous bases are held together by hydrogen bonds, which maintain the shape of the DNA ladder. These base pairs are formed when a pyrimidine binds with a purine. Specifically, adenine forms a base pair only with thymine via two hydrogen bonds, and cytosine forms a base pair with only guanine via three hydrogen bonds (Figure 2.1). A skew and a twist of the strands of DNA lets in as little water as possible between the nitrogenous bases, forming the well-known double helix shape of the double stranded DNA (dsDNA) molecule.

Graphical user interface, application

Description automatically generated with medium confidence**Cellular DNA**

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**Nucleus**

***Wolbachia***

**Bacterial chromosome**

***Phage WO DNA***

**Mitochondria**

**Mitochondrial DNA**

***Phage WO DNA***

**Eukaryotic chromosomes**

**Figure 2.2.** A eukaryotic cell containing nuclear DNA (gold), mitochondrial DNA (teal), bacterial DNA (blue) and bacteriophage DNA (red).

Arthropod cells may contain multiple types of DNA (Figure 2.2).

1. *Nuclear DNA*: Eukaryotic DNA is coiled and condensed around nuclear proteins, called histones, to form chromosomes. Chromosomes are stored in the nucleus of each cell and encode the arthropod’s genetic information.
2. *Mitochondrial DNA*: Mitochondria, the organelle responsible for cellular respiration and energy production, contains its own DNA. Mitochondrial DNA is circular and, like its encoding organelle, is passed down from mother to offspring.
3. *Bacterial DNA: Wolbachia* and other bacterial endosymbionts contain a single, circular, supercoiled DNA molecule called the bacterial chromosome.
4. *Bacteriophage DNA:* Phage WO, the bacteriophage that infects *Wolbachia,* carries its own genome. Phage WO is a temperate phage, meaning it can integrate its genome into the bacterial chromosome. When active, WO packages its DNA into phage capsids and lyses the *Wolbachia* cell to infect new hosts. Therefore, phage DNA can be found in both the *Wolbachia* chromosome and in phage particles.

In addition to the DNA shown above, an arthropod cell may also contain eukaryotic viruses and other intracellular bacteria (e.g., *Rickettsia*, *Spiroplasma*, *Cardinium*).

*Graphical user interface, application

Description automatically generated with medium confidence****Wolbachia* Localization**

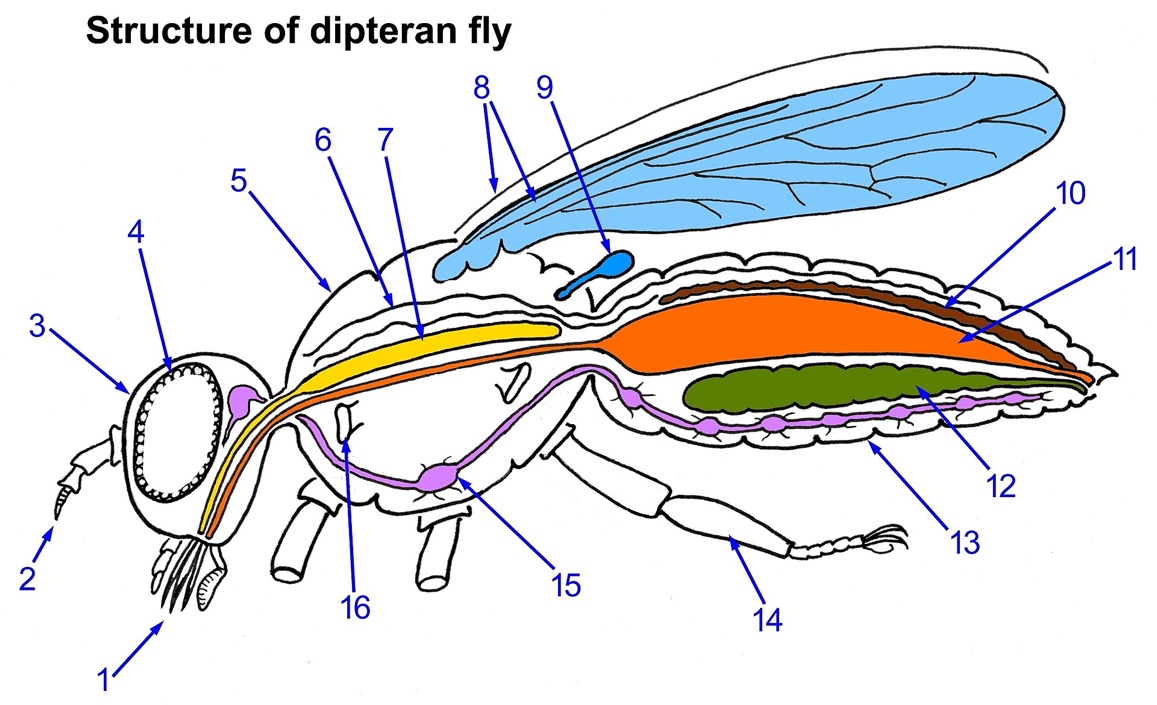
https://theminione.com/minione-resource-center/

In most cases, whole body DNA extractions are sufficient to obtain *Wolbachia* DNA. If a specimen is particularly large and/or has a tough exoskeleton, abdominal dissections are recommended. A general rule of thumb is to dissect specimens that are larger than a grain of rice, such as a fruit fly, small ant, or mosquito. Because *Wolbachia* is maternally transmitted, it is commonly localized in the reproductive organs (ovaries and testes); however, increasing studies have also detected the symbiont in somatic, non-reproductive tissue. For the purpose of this lab activity, we recommend dissecting the reproductive areas. Make sure the arthropod is not still alive; freezing overnight and/or preserving in alcohol should be done prior to dissections and DNA extraction.

**General guidelines for *Wolbachia* DNA extractions:**

1. *Small arthropods:* If the specimen is about the size of a grain of rice (such as a fruit fly or small ant ~2 mm), extract DNA from the entire body.
2. *Large arthropods:* If the specimen is larger than ~2 mm, use a scalpel or razor blade to cut off the abdomen. In many cases, such as grasshoppers and cockroaches, the abdomen may still be larger than 2 mm. If so, use the scalpel to extract an internal portion of the abdomen most likely to contain reproductive tissue. The anatomy of a Dipteran fly is shown below for reference (Figure 2.3). Research the anatomy of each arthropod prior to dissections.
3. *Tough arthropods:* If the specimen has a particularly tough exoskeleton, such as a tick, use a scalpel or razor blade to carefully cut open the abdomen and expose internal organs. This will ensure that the cell lysis buffer has access to reproductive tissues.

*Image by Daktaridudu; available in Wikipedia under CC BY-SA 4.0 license*



**Paired antennae**

**Complex mouthparts**

**Head**

**Compound eye**

**Thorax**

**Open heart tube**

**Salivary gland**

**Wings**

**Halter**

**Excretory (Malpighian) tube**

**Gut**

**Ovary (or Testis)\***

**Abdomen**

**Legs**

**Thoracic ganglion of nervous system**

**Spiracle opening to respiratory tube (trachea)**

***\* Wolbachia-enriched***

**Figure 2.3.** Dipteran fly anatomy

Graphical user interface, application

Description automatically generated with medium confidence**Technical Overview**

https://theminione.com/minione-resource-center/

This DNA extraction protocol is based on the [insect adaptation](https://www.qiagen.com/us/resources/resourcedetail?id=cabd47a4-cb5a-4327-b10d-d90b8542421e&lang=en) of Qiagen’s DNeasy Blood & Tissue kit (Product # 69504).

**Sample Preparation**

Each specimen will be rinsed with water to remove alcohol preservative. If larger than a fruit fly, the abdomen will be dissected to obtain the reproductive tissues. If the specimen contains a particularly tough exoskeleton, abdominal tissue can be exposed using a scalpel or razor blade.

**Cell Lysis & DNA Precipitation**

Each specimen will be macerated in a cell lysis solution to break open the cellular and nuclear membranes. As a result, DNA is exposed to proteases, such as nucleases, in the host tissue. Therefore, the enzyme *Proteinase K* must be added to denature the proteins and keep the DNA intact. The solution is heated to ~ 56°C for enhanced lysis. Finally, ethanol will be added to bring the DNA out of solution.

**DNA Purification**

Once the proteases are destroyed and DNA is precipitated, the DNA must be purified. All cellular components, including DNA, will be placed into a spin column. Upon centrifugation, the material will pass through the filter, which attracts DNA but allows cellular debris to pass through. This will be followed by two wash steps with two separate buffers, AW1 and AW2.

***DNA Elution***

The activity will be completed by removing the DNA from the filter. This is called *eluting the DNA* and is done by adding the elution Buffer AE. DNA is more attracted to the elution buffer than the filter, and finally passes through the filter. Spinning the tube with the DNA embedded in the filter will pull the elution buffer through the matrix, thus collecting the DNA into the 1.5 ml tube.

Graphical user interface, application

Description automatically generated with medium confidence**Pre-Lab Questions**

https://theminione.com/minione-resource-center/

**Read through the entire lab activity and answer the questions below.**

1. Based on your knowledge of DNA structure, what is the complementary strand of this DNA sequence?

**5’ – ATG CCG GAA TCG TTA GCA – 3’**

1. Is it possible for your DNA extraction to contain only *Wolbachia* DNA? Explain why or why not.
2. List the two controls used in this experiment. Why do we need controls at this stage?
3. Which centrifuge rotor(s) is properly balanced?

1

2

3

4

5

6

7

8

7

6

5

4

3

1

5

4

1

C

B

A

2

8

2

8

7

3

6

1. How would you properly balance the other rotor(s)?
2. What would happen if you used the same pipette tip throughout the entire protocol?

Graphical user interface, application

Description automatically generated with medium confidence**Pre-Lab Questions, Part 2**

https://theminione.com/minione-resource-center/

1. Based on background research, **formulate a hypothesis** about the frequency of *Wolbachia* endosymbionts in your specimens.
2. What is the purpose of each reagent?

Proteinase K:

Buffer AL:

Ethanol:

Buffer AW1:

Buffer AW2:

Buffer AE:

1. Which centrifugation step will transfer DNA to the spin column membrane (or filter)?
2. Which centrifugation step will remove, or elute, DNA from the spin column?
3. Indicate which pipette (P200 or P1000) should be used for each step.

Step 5:

Step 8:

Step 9:

Step 12:

Step 14:

Step 16:

Step 19:

Step 21:

Graphical user interface, application

Description automatically generated with medium confidence**DNA Extraction Protocol**

https://theminione.com/minione-resource-center/

**Class Materials**

**Materials per Group**

*\* Only needed if using water bath*

* Mini-centrifuge
* Float rack \*
* Metal tongs \*
* Incubator, heat block, or water bath, 56 °C
* Vortex mixer

*\* Only needed if using water bath*

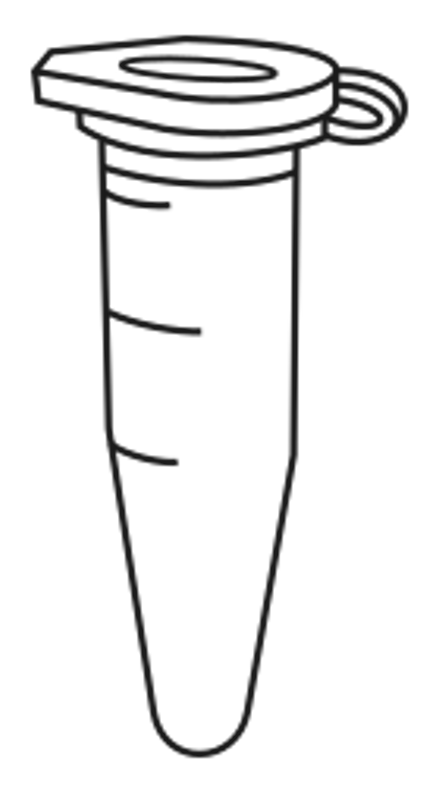
* 2 Arthropod specimens
* +/- *Drosophila* controls
* Gloves
* Tweezers / Scalpel
* Petri dish
* Water
* Transfer pipette or squirt bottle
* 4 Microtube pestles
* 12-1.5 ml tubes
* Qiagen DNeasy Kit aliquots
* Buffer ATL (900 μl)
* Proteinase K (100 μl)
* Buffer AL (1.0 ml)
* Buffer AW1 (2 x 1.25 ml)
* Buffer AW2 (2 x 1.25 ml)
* Buffer AE (500 μl)
* Ethanol (95-100%, 1.0 ml)
* 4 spin columns
* 4 collection tubes
* P200 and P1000 pipettes
* P200 and P1000 tips
* Waste cup for tips
* Waste cup for liquids
* Sharpie
* Kimwipes
* Tube rack

***Sample Preparation***

1. Label the contents of each tube in the chart below.

|  |  |  |
| --- | --- | --- |
| Group #  (or initials) | Tube # | Contents |
|  | 1 |  |
|  | 2 |  |
|  | 3 | Positive (+) Arthropod Control |
|  | 4 | Negative (-) Arthropod Control |

1. If preserved in alcohol, use tweezers to carefully transfer the first arthropod to a Petri dish. Rinse with water using either a transfer pipette or squirt bottle and blot dry excess liquid. Take 1-2 pictures for the [The Wolbachia Project Database.](https://wolbachiaprojectdb.org/) If the arthropod is smaller than a grain of rice, skip to Step #4.
2. Remove the abdomen of the arthropod and cut off a small piece (roughly ~2 mm, or small enough to fit in the bottom of a microcentrifuge tube).



**1.5 mL microcentrifuge tube**

*Remove as much preservative and water as possible. If the arthropod is large or has a thick/tough exoskeleton, dissect out the reproductive tissues. Arthropods with a thick exoskeleton should be cut into multiple pieces.*

1. Place the specimen into a labeled 1.5 ml microcentrifuge tube and repeat for remaining arthropods.

***Cell Lysis & DNA Precipitation***

1. Add 180 μl Buffer ATL to the first tube.

*Buffer ATL is a tissue lysis buffer.*

1. Use a sterile pestle to grind the sample for 1 minute.

*This is the most critical step of the entire protocol. Thorough grinding is necessary to obtain high DNA yield. Use muscle power and grind each sample thoroughly.*

1. Using a new pestle for each tube; repeat Steps 5 and 6 with remaining samples.
2. Add 20 μl Proteinase K to each tube.

*Proteinase K will destroy DNases that break down DNA.*

1. Add 200 μl Buffer AL to each tube and immediately mix by vortexing for 10 seconds or pipetting up and down.

*Buffer AL lyses open cells.*

1. Incubate for at least 15 minutes @ 56 °C. Longer incubation times (i.e., 2-3 hours) are most effective for cellular lysis.
2. Centrifuge (spin) the tubes for about 30 seconds to pellet debris as this could clog the filter of the spin column. Using a new tip for each sample, transfer the supernatant (~400 μl) to a labeled 1.5 ml tube. Discard the old tube of cellular debris and repeat for other samples.

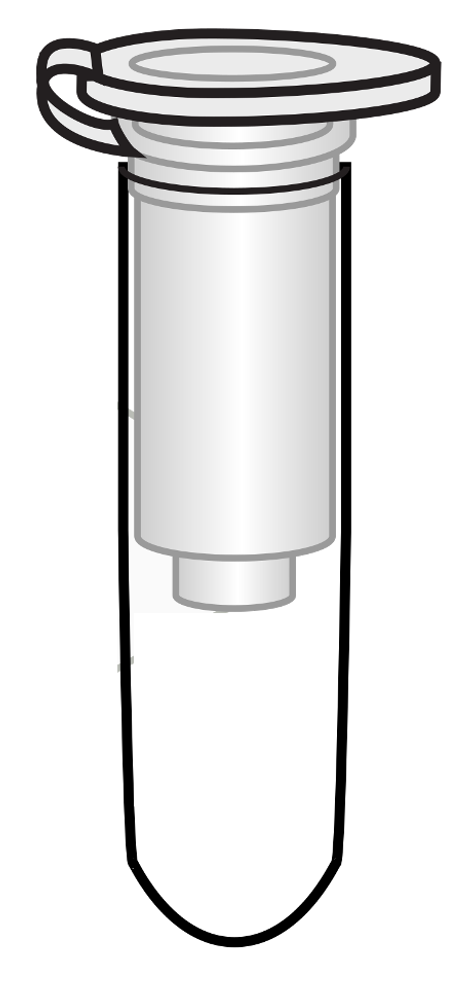
*Use caution to not disturb the pellet. Alternatively, pour the DNA-containing supernatant directly from each sample into the corresponding 1.5 ml tube.*

1. Add 200 μl ethanol (96-100%) to each tube and mix by vortexing for 10 seconds or pipetting up and down.

*Ethanol precipitates DNA.*

**This is an optional STOPPING POINT. Store DNA in in refrigerator (4 °C) until next class period.**

***DNA Purification***



**Spin column**

**2.0 mL collection**

**tube**

1. Collect four DNeasy spin columns fitted with four 2.0 ml collection tubes and label the lids of the spin columns with your sample numbers.
2. Pipet (or carefully pour) the liquid from Tube 1 of the above steps (including any precipitate) into DNeasy Mini column #1. Using a new pipette tip for each transfer, repeat this process with the three other tubes. Make sure to keep the tube numbers consistent.

*Throughout the protocol, be careful not to touch the filter or get liquid around the rim of the column.*

1. Centrifuge all tubes for 1 minute at ≥6,000 x g (8,000 rpm).The DNA is now caught in the filter of the spin column. Carefully remove the spin columns from the collection tubes and discard the flow through (liquid) from the 2.0 ml collection tubes into the waste cup. Replace the spin columns back into their emptied collection tubes.

*If using a mini-centrifuge, you may not be able to fit all columns in the rotor. Spin 2-3 at once, ensuring that the centrifuge is properly balanced at all times. If your centrifuge does not have an adjustable speed, use the default speed throughout the protocol.*

1. Add 500 μl of Buffer AW1 to each spin column.

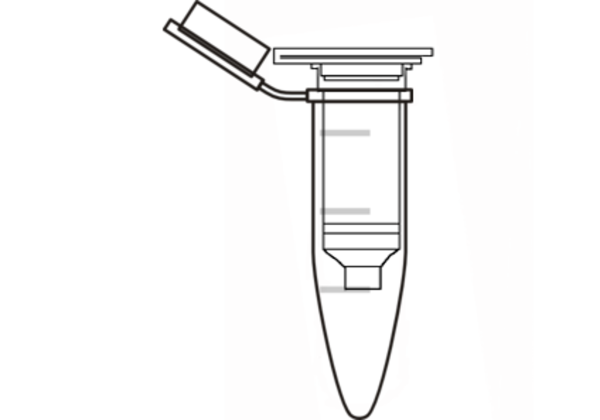
*Buffer AW1 washes the DNA.*

1. Centrifuge all tubes for 1 minute at ≥6,000 x g (8,000 rpm).
2. Again, discard the flow through waste from the 2.0 ml collection tubes into the waste cup and place the DNeasy Mini Spin Columns back into the same 2.0 ml collection tubes.
3. Add 500 μl Buffer AW2 (a second wash buffer) to each of the four tubes and centrifuge for 3 minutes at 20,000 x g (14,000 rpm), or max speed of your centrifuge if it doesn’t go that high, to dry the DNeasy membrane.

*If the centrifuge you are using cannot attain this speed, allow the tube to air dry for 5 minutes. This will evaporate the ethanol. It is important to air dry the membrane so that ethanol does not interfere with PCR.*

***DNA Elution***

1. Transfer the spin columns into 1.5 ml microcentrifuge tubes. Be sure to label the lids of each tube #1-4 and include your group number (or initials). These will contain the purified DNA samples.



**Spin column**

**1.5 mL**

**microcentrifuge**

**tube**

*Carefully remove the DNeasy column so that it does not make contact with residual ethanol in the tube. If ethanol is still present on the membrane, empty the collection tube and spin again.*

*At this point, you will have two sets of lids. When you place them in the rotor, rotate the column such that its lid is not directly on top of the 1.5 ml tube lid. Keep lids down against the center of the rotor, not sticking out, and face them in the direction of motion. If a lid breaks off, transfer DNA to a new labeled 1.5 ml tube.*

1. Pipet 100 μl of Buffer AE directly onto the spin column membrane. Incubate at room temperature for 1 minute.

*Buffer AE is is an elution buffer that rinses the DNA off the spin column filter and into the 1.5 ml tube. To incubate at room temperature, simply leave the tube sitting on the lab bench for 1 minute.*

1. Centrifuge at ≥6,000 g or 8,000 rpm for 1 minute. **The eluted DNA will be collected in the 1.5 ml tube.**
2. Discard the spin columns and KEEP the labeled 1.5 ml tubes. The DNA is now in the 1.5 ml microcentrifuge tubes. *Optional:* Incubate the DNA for 1 hour @ 65 °C or overnight at room temperature.
3. Store the eluted DNA frozen at -20 °C until PCR.

Graphical user interface, application

Description automatically generated with medium confidence**DNA Extraction Bench Protocol**

https://theminione.com/minione-resource-center/

*This is an abbreviated protocol. Make sure to label all tubes and change pipette tips between samples.*

***Sample Preparation***

* + Remove the abdomen of the arthropod and cut off a small piece (roughly ~2 mm, or small enough to fit in the bottom of a microcentrifuge tube). If the specimen is smaller than a grain of rice, use the entire body.
  + Place the specimen in a labeled 1.5 ml microcentrifuge tube.

***Cell Lysis & DNA Precipitation***

* + Add 180 μl Buffer ATL and use a sterile pestle to grind the sample for 1 minute.
  + Add 20 μl Proteinase K.
  + Add 200 μl Buffer AL and immediately mix by vortexing for 10 seconds or pipetting up and down.
  + Incubate for *at least* 15 minutes @ 56 °C. Longer incubation times (i.e., 2-3 hours) are most effective for cellular lysis.
  + Spin the tube for about 30 seconds to pellet debris. Carefully transfer the supernatant to a new labeled 1.5 ml microcentrifuge tube. Discard the tube of cellular debris.
  + Add 200 μl ethanol (96-100%) and mix by vortexing for 10 seconds or pipetting up and down.

Optional stopping point: store DNA in refrigerator (4 °C)

***DNA Purification***

* + Label a DNeasy spin column fitted with a 2.0 ml collection tube.
  + Pipet the liquid containing ethanol-precipitated DNA into the DNeasy spin column.
  + Centrifuge for 1 minute at ≥6,000 x g (8,000 rpm).Discard the flow through from the 2.0 ml collection tube.
  + Add 500 μl of Buffer AW1 and centrifuge for 1 minute at ≥6,000 x g (8,000 rpm). Discard the flow through.
  + Add 500 μl Buffer AW2 and centrifuge for 3 minutes at 20,000 x g (14,000 rpm).

***DNA Elution***

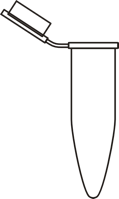
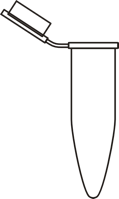
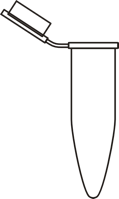
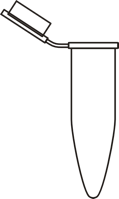
* + Transfer the spin columns to a labeled 1.5 ml microcentrifuge tube. Discard the 2.0 ml collection tube.
  + Pipet 100 μl of Buffer AE directly onto the spin column membrane.
  + Incubate at room temperature for 1 minute.
  + Centrifuge at ≥6,000 g or 8,000 rpm for 1 minute.
  + Discard the spin column and KEEP the labeled 1.5 ml tube.
  + *Optional:* incubate the DNA for 1 hour @ 65 °C or overnight at room temperature.
  + Store the eluted DNA frozen at -20 °C until PCR.

Graphical user interface, application

Description automatically generated with medium confidence**Post-Lab Questions**

https://theminione.com/minione-resource-center/

1. Illustrate the contents of each tube upon completion of the following steps. Below, label where the DNA is located.



Sample Preparation

Cell Lysis

Cellular Debris Removal

DNA Elution

1. After completion of the DNA extraction, what did the DNA look like?
2. Where is *Wolbachia* generally located within an arthropod? Why?
3. What is the purpose of isolating the DNA? What is your next step in determining the frequency of *Wolbachia*?
4. Imagine that there is an *E. coli* outbreak in your state and you would like to test the lettuce from your local grocery store. How could you modify this protocol in order to extract DNA from the lettuce (to identify the species) and check for presence/absence of *E. coli*. ? Keep in mind that (i) *E. coli* is free-living and not an endosymbiont, and (ii) plant cells are encased in both a cell membrane *and* cell wall.

Graphical user interface, application

Description automatically generated with medium confidence**Database Entry**

https://theminione.com/minione-resource-center/

After completing the DNA Extraction, open your entries in The *Wolbachia* Project Database and record Methods and protocol notes. A comprehensive guide is located under the Resources tab.

[**https://wolbachiaprojectdb.org/**](https://wolbachiaprojectdb.org/)

**Database Fields to Complete**

**Methods**

* DNA extraction kit
* DNA extraction location
* Update protocol notes