**HEART RATE RESPONSE TO INDUCED STIMULI IN FRESHWATER SHRIMP**

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**Purpose**

To investigate the effect of various environmental cues upon the heart rate of the transparent ghost shrimp, *Palaemonetes kadakensis*. There are many factors that affect the heart rate response including temperature and chemical stimulants or depressants. For the purposes of this experiment, cold water and nicotine will serve as the temperature and chemical stimuli.

**Preparation**

The ghost shrimp is an ideal experimental model for monitoring heart rate, due to the organism’s transparent exoskeleton and low maintenance. In addition, the animals are easily acquired, making them well suited for use in the classroom. Before starting this experiment, become familiar with the shrimp anatomy terms below:

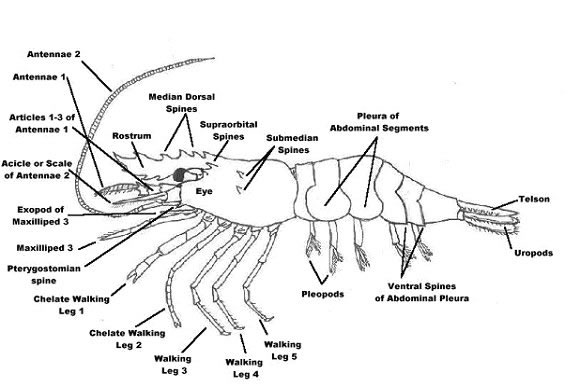


FIG 1.*General shrimp anatomy diagram. Source: <http://www.geog.ubc.ca/biodiversity/efauna/*

*CaridaeShrimpsandPrawnsofBC.html>*

**Introduction**

Heart rate in crustaceans can be altered under many conditions. Neurotransmitters, temperature, and chemicals such as stimulants can all have an effect. Neurotransmitters act on the organism’s heart rate through the nervous system in a parasympathetic-like or sympathetic-like manner. This can either cause an increase or decrease in overall heart rate based on the properties of the neurotransmitter in question. The effects of temperature on heart rate are also variable. Lower temperatures tend to decrease the heart rate. Conversely, high temperatures tend to cause an increase in heart rate due to the increase in metabolic activity and higher rate of chemical reactions within the body. Chemical stimulants also increase the heart rate and blood flow. In this experiment, students will become familiar with these effects by subjecting a species of freshwater shrimp to varying environmental conditions.

**Materials**

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| Item | Quantity |
| Ghost Shrimp (*Palaemonetes kadakensis*) | 1 |
| Small Wooden Rod (Toothpick) | 1 |
| Grooved Petri Dish | 1 |
| Light Microscope | 1 |
| Pins with Curled Ends | 2 |
| Paper Towels | 2 |
| Beaker of Distilled, Aerated Water | 2 |
| Beaker of Ice | 1 |
| 10 μM Nicotine Solution | 1 Bath |
| 10 μM GABA Solution | 1 Bath |
| 10 μM Glutamate Solution | 1 Bath |
| Set of Microscope Lights | 1 |
| Drop of Super Glue (Maxi-Cure) | 1 |
| Drop of Quick Dry (Insta-Set) | 1 |

FIG 2.*Materials for shrimp preparation.*

**Methods**

1. Dampen a paper towel with distilled water as this species of freshwater shrimp are highly sensitive to small amounts of copper or nickel.
2. Place the shrimp in the damp paper towel so that the anterior region is wrapped and the animal’s back and tail are free.
3. With a second, dry paper towel gently dab the back of the animal in order to dry it for the glue.
4. Place a small drop of the glue onto the wooden rod and then adhere it to the animal. Ensure that the stick is glued slightly below the back on the tail so that the heart is not hidden by the apparatus.
5. While holding the rod in place, place a dab of the quick dry compound over the glue to complete the bonding.
6. Hold the stick in place for approximately five more seconds or until the rod is sufficiently affixed to the shrimp.

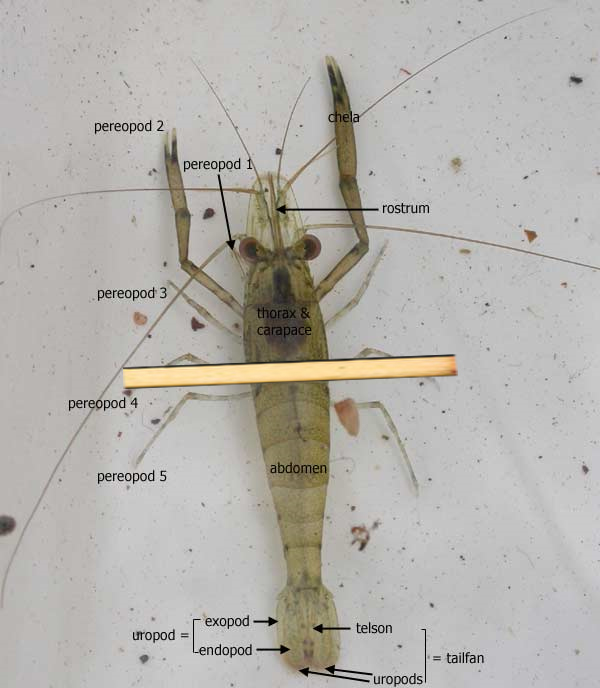


FIG 3.*Diagram of wooden rod placement on dorsal side of shrimp. Source: <www.mdfrc.org.au/BugGuide/diagrams/decapoda.htm*>

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1. Rinse the shrimp and rod of any excess chemicals by dipping it once or twice into the beaker of distilled water.
2. Place the shrimp in the grooved Petri dish; ensure that the rod fits securely into both grooves on either side of the dish. This will limit the animal’s movement and will allow for it to be monitored through the microscope. Fill the dish with aerated water and be sure that the level is over the back of the shrimp so that it survives the experiment.

**Exercise 1.**

Monitor the heart rate of the shrimp to gain a baseline. To do this, place the shrimp setup under the microscope so that the heart is visible on the back of the animal. Count the number of beats in ten seconds and multiply this number by six to get the beats per minute. To obtain an experimental baseline, do this three times and take the average of the three values. Note: It may be beneficial to allow the shrimp to acclimate for approximately five to ten minutes before taking a baseline reading to account for the agitation of the animal.

Heart Rate

1. \_\_\_\_\_\_\_\_\_\_beats/minute
2. \_\_\_\_\_\_\_\_\_\_beats/minute
3. \_\_\_\_\_\_\_\_\_\_beats/minute

\_\_\_\_\_\_\_\_\_\_average beats/minute (BASELINE)

**Exercise 2.**

**Note:** Your TA will instruct your group to perform either Part A. or Part B. of Exercise 2 (NOT both). You will then exchange recorded data with a group that used a different chemical stimulus.

**A.** Carefully transport the dish containing the shrimp to the sink and gently pour the contents of the bath out. Make sure that the shrimp is secure and does not fall into the sink. While monitoring the shrimp heart rate through the microscope, fill the bath with the prepared glutamate solution. Take note of the immediate response. After approximately thirty seconds, take note of the number of heart beats in a ten second period. Again, multiply this number by six in order to calculate beats per minute. As in Exercise 1, repeat this process for a total of three times; generate an average beats per minute reading.

Heart Rate

1. \_\_\_\_\_\_\_\_\_\_beats/minute
2. \_\_\_\_\_\_\_\_\_\_beats/minute
3. \_\_\_\_\_\_\_\_\_\_beats/minute

\_\_\_\_\_\_\_\_\_\_average beats/minute

**B.** Carefully transport the dish containing the shrimp to the sink and gently pour the contents of the bath out. Make sure that the shrimp is secure and does not fall into the sink. While monitoring the shrimp heart rate through the microscope, fill the bath with the prepared GABA solution. Take note of the immediate response. After approximately thirty seconds, take note of the number of heart beats in a ten second period. Again, multiply this number by six in order to calculate beats per minute. As in Exercise 1, repeat this process for a total of three times; generate an average beats per minute reading.

Heart Rate

1. \_\_\_\_\_\_\_\_\_\_beats/minute
2. \_\_\_\_\_\_\_\_\_\_beats/minute
3. \_\_\_\_\_\_\_\_\_\_beats/minute

\_\_\_\_\_\_\_\_\_\_average beats/minute

**Exercise 3.**

Carefully transport the dish containing the shrimp to the collection beaker in the classroom. Gently pour out the solution containing either GABA or glutamate. Make sure that the shrimp is secure. Rinse the beaker and the shrimp with aerated water to ensure that all of the chemicals have been removed. While monitoring the shrimp heart rate through the microscope, fill the bath with chilled aerated water from the beaker containing the ice. Observe the immediate change. After approximately thirty seconds, take note of the number of heart beats in a ten second period. Again, multiply this number by six in order to calculate beats per minute. As in the previous exercises, repeat this process for a total of three times; generate an average beats per minute reading.

Heart Rate

1. \_\_\_\_\_\_\_\_\_\_beats/minute
2. \_\_\_\_\_\_\_\_\_\_beats/minute
3. \_\_\_\_\_\_\_\_\_\_beats/minute

\_\_\_\_\_\_\_\_\_\_average beats/minute

**Exercise 4.**

Nicotine will be applied to the bath last, as it is expected induce a significantly stronger change in heart rate than the stimuli used in Exercises 2 and 3. In order to isolate the effects of nicotine from experimental stress, a chilled solution of nicotine will be used. Carefully transport the dish containing the shrimp to the sink and gently pour the contents of the bath out. Make sure that the shrimp is secure and does not fall into the sink. While monitoring the shrimp heart rate through the microscope, fill the bath with chilled nicotine solution from the beaker indicated by the TA. Observe the immediate change. After approximately thirty seconds, take note of the number of heart beats in a ten second period. Again, multiply this number by six in order to calculate beats per minute. As in the previous exercises, repeat this process for a total of three times; generate an average beats per minute reading.

Heart Rate

1. \_\_\_\_\_\_\_\_\_\_beats/minute
2. \_\_\_\_\_\_\_\_\_\_beats/minute
3. \_\_\_\_\_\_\_\_\_\_beats/minute

\_\_\_\_\_\_\_\_\_\_average beats/minute

**Exercise 5.**

In order to compare the effects upon heart rate of the various conditions observed above, graph the average beats/minute of each stimulus below:

**Heart Rate (Beats/Min)**

**Control GABA Glutamate Cold Nicotine**

**Exercise 6.**

To account for the effects of non-target variable conditions such as experimenter noise or changes in the amount of light over the shrimp with passing shadows, it is important to calculate the percent difference from baseline for each of the stimuli observed before drawing any conclusions about generalized reaction trends. The formula for percent difference is as follows:

For the conditions GABA, glutamate, and cold water, the baseline used will be the average beats/min for the room temperature water (control) from Exercise 1. For the nicotine trial, the baseline used will be the average beats/min for the chilled water from Exercise 3.

Percent Differences:

GABA: \_\_\_\_\_\_\_\_\_\_\_\_\_

Glutamate: \_\_\_\_\_\_\_\_\_\_\_

Cold Water: \_\_\_\_\_\_\_\_\_\_

Nicotine: \_\_\_\_\_\_\_\_\_\_\_\_

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