	WW.	Environmental Analysis Teaching	Date: 8/10/2016	Number: 07B v.03
	POMONA	and Research Laboratory		
		Standard Operating Procedure	Title: Using Micro-Pippettes	
	POMONA COLLEGE	Approved By: TBD	Revision Date: 5/10/23	

1. Scope and Application

- 1.1 The scope of this SOP is to train researchers to learn and follow the basic protocol of using, cleaning, and operating everyday lab equipment, which includes, the micropipette, glassware equipment, measuring balances, the Vortex, and other marginal lab tools that will be further specified.
- 1.2 The applications of this SOP are for any basic lab setting for students conducting experiments and/or learning new lab procedures.

2. Summary of Method

2.1

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- 3. Acknowledgements
- 4. Definitions
 - **4.1** Term1:
- 5. Health and Safety
 - **5.1** Describe the risk...

Safety and Personnnel Protective Equipment

- 6. Personnel & Training Responsibilities
 - **6.1** Researchers training is required before this the procedures in this method can be used...
 - **6.2** Researchers using this SOP should be trained for the following SOPs:
 - SOP01 Laboratory Safety
- 7. Apparati
 - 7.1 Pipetman (ranging from 2-1000microliters)

8. What is a Micropipette?

Micropipettes are used to transfer small amounts (i 1 ml) of liquids. The scales on micropipettes are in microliters (1000 l = 1 ml). These are very expensive, delicate instruments so be very careful during use. The sizes of the micropipettes can be identified via the number on the round button on the plunger; this value is the maximum volume in microliters that can be transferred with that size pipette. The following is an illustration of a micropipette:

9. How to Use a Micropipette

1. NEVER exceed the upper or lower limits of these pipettes. The limits are: P10: 1.0 - 10.0 l; P20: 2.0 - 20.0 l; P200: 20 - 200 l; P1000: 200 - 1000 l Look at the front face of the pipet and you will see a window with three digits inside. The diagram below shows the MAXIMUM value that can or should be dialed in on each size pipet. To exceed these values will put the pipet out of calibration. Beside each "window" below is the numbers place it represents. (figure 2)

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Figure 1: The Pipetman

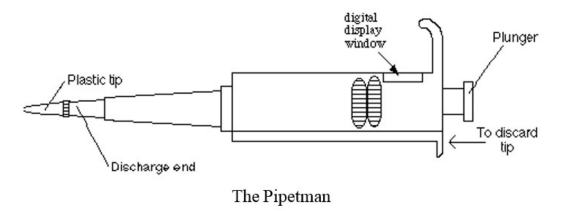
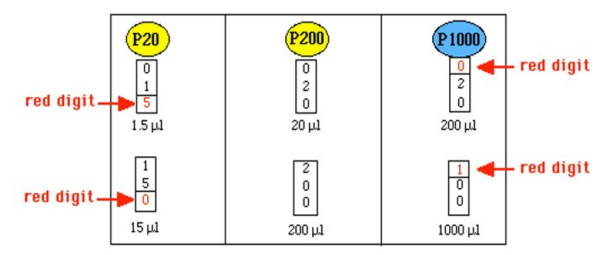


Figure 2: Pipetman Window



- RULE OF THUMB: Always select the SMALLEST size pipette that will handle the volume you wish to move to achieve the greatest accuracy. Accuracy decreases as you use unnecessarily large pipets for small volumes.
- 2. Set the desired volume by turning the centrally located rings clockwise to increase volume or counterclockwise to decrease volume.
- 3. Load a sterile tip. Use the appropriate tips for each micropipette as designated by the labels of the box, and by confirming with the lab instructor.
- 4. 5. Load the sample.
 - The plunger will stop at two different positions when it is depressed. Push the plunger down slowly to the point of first resistance: this is the load volume. Because this first stopping point is dependent on the volume that is being trans-

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ferred, the distance you have to push the plunger to reach the point of initial resistance will change depending on the volume being pipetted.

- While holding the plunger at the load volume set point, put the tip into the solution so that it is immersed just enough to cover the end (3-4 mm), not as deep as possible.
- Slowly release the plunger to draw up the liquid making sure to keep the tip immersed. NOTE: If the solution you are pipetting is viscous, allow the pipet tip to fill to final volume before removing it from solution to avoid the presence of bubbles in the plastic tip, which will result in an inaccurate volume.
- Visually inspect the load to make sure it is correct there should be no air space in the distal end tip.

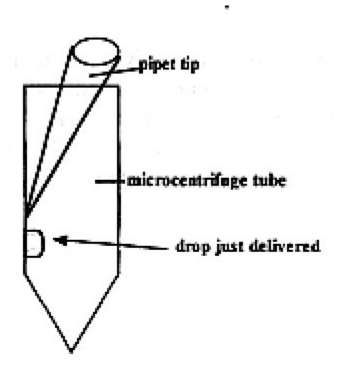


Figure 3: Loading Sample

- 5. Deliver the sample. The second stopping point can be found when the plunger is depressed beyond the initial resistance until it is in contact with the body of the pipette. This second stopping point is used for the complete discharging of solutions from the plastic tip. You should not reach this second stop when drawing liquid into the pipette, only when expelling the last drop.
- 6. Discharge the tip. While holding the tip over an appropriate waste receptacle, press the discharge slider on the back of the grip.

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10. Things to Keep in Mind

 Do not hesitate to ask the lab instructor questions about the pipetting protocol and measures to avoid contamination

- Never point a pipette up. This may cause liquid to run down into the pipette destroying it
- When withdrawing liquids with the pipette, always release the plunger slowly. This prevents liquid from rushing into the end of the pipette and clogging it up. This is especially important with large volume pipettes
- Be sure you use the proper size tip for each pipette.
- Always use a new tip for each different liquid.
- Use the correct pipette for the volume that is to be dispensed. Going below or above the range of the micropipette may damage the instrument.

11. QC/QA Criteria

12. References

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