Titration of Aspirin Tablets

In this lab, you will determine the percent purity of two commercially available aspiring tablets using an acid-base titration. In general, an acid and a base react to produce a salt and water by transferring a proton (H^+) :

$$\begin{array}{rcl} \text{HA}(aq) + & \text{NaOH}(aq) \rightarrow & \text{H}_2\text{O}(l) + & \text{NaA}(aq) \\ \text{acid} & & \text{base} & & \text{salt} \end{array}$$
(1)

The active ingredient in aspirin, and the chemical for which aspirin is the common name, is acetylsalicylic acid. To determine the amount of aspirin (acetylsalicylic acid) in a sample, the precise volume and concentration of the NaOH, and the overall reaction, must be known. The NaOH serves as a secondary standard, because its concentration can change over time. To find the precise concentration of the NaOH, it must be titrated against a primary standard, an acid that dissolves completely in water, has a high molar mass, that remains pure upon standing, and is not hygroscopic (tending to attract water from the air). Because sodium hydroxide is hygroscopic, it draws water from its surroundings. This mean one cannot simply weigh out a sample of sodium hydroxide, dissolve it in water, and determine the number of moles of sodium hydroxide present using the mass recorded, since any sample of sodium hydroxide is likely to be a mixture of sodium hydroxide and water. Thus, the most common way to determine the concentration of any sodium hydroxide solution is by titration. Determining the precise concentration of NaOH using a primary standard is called standardization. You will first standardize your NaOH solution, and then use it to analyze aspirin tablets for their aspirin content and purity.

An aspirin tablet may also include inactive ingredients that help produce a consistent product for consumers. Think of the purpose these ingredients serve when considering whether your percent purity is reasonable for a commercial aspirin tablet.

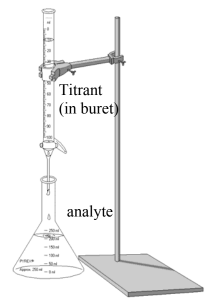
What is a titration?

A titration is a procedure for determining the concentration of a solution (the analyte) by allowing a carefully measured volume of this solution to react with another solution whose concentration is known (the titrant). The point in the titration where enough of the titrant has been added to react exactly with the analyte is called the equivalence point, and occurs when moles of titrant equals moles of analyte according to the balanced equation. For example, if a monoprotic acid (the analyte) is titrated with a strong base like sodium hydroxide (the titrant), the equivalence point occurs when

number of moles of
$$OH =$$
 number of moles of HA. (2)

The equivalence point is often marked by an indicator, a substance that changes color at (or very near) the equivalence point.¹

There are many types of titrations. In this lab you will be performing an acid base titration.



¹ The point at which the indicator changes color is called the "endpoint". We will normally assume that the endpoint is equal to the equivalence point.

Background

Standardizing a Base

Consider the following reaction between oxalic acid, $H_2C_2O_4(aq)$, and potassium hydroxide:

$$H_2C_2O_4(aq) + 2 \operatorname{KOH}(aq) \rightarrow 2 H_2O(l) + K_2C_2O_4(aq)$$
(4)

If 0.4862 g of oxalic acid was dissolved in water and titrated with 17.98 mL of potassium hydroxide solution, the concentration of the potassium hydroxide solution can be calculated. Since 17.98 mL of potassium hydroxide solution is used, that volume is converted to liters and put into the denominator:

molarity of KOH = [KOH] =
$$\frac{\text{mol KOH}}{0.01798 \text{ L KOH}}$$
 (5)

The mass of oxalic acid, $H_2C_2O_4(aq)$, and Equation (4) are then used to determine the number of moles of potassium hydroxide present:

$$0.486 \text{ g } \text{H}_2\text{C}_2\text{O}_4 \times \frac{\text{mol } \text{H}_2\text{C}_2\text{O}_4}{90.036 \text{ g } \text{H}_2\text{C}_2\text{O}_4} \times \frac{2 \text{ mol } \text{KOH}}{1 \text{ mol } \text{H}_2\text{C}_2\text{O}_4} = 0.0108 \text{ mol } \text{KOH} \quad (6)$$

Finally, the molarity for potassium hydroxide is calculated as follows:

molarity of KOH = [KOH] =
$$\frac{0.01080 \text{ mol KOH}}{0.01798 \text{ L KOH}} = 0.6007 \text{ M KOH}$$
 (7)

Determining the Number of Moles of an Acid

Once a solution has been standardized, the solution can be used to determine the molar concentration of another solution, or simply the number of moles of analyte in the flask. Combined with the precise molar concentration of the titrant, the precise volume of titrant delivered yields the number of moles used to react with the analyte. Consider the following reaction between a monoprotic acid and potassium hydroxide:

$$HC_{9}H_{7}O_{4}(s) + KOH(aq) \rightarrow H_{2}O(l) + KC_{9}H_{7}O_{4}(aq)$$
(8)

Suppose 0.375 grams of acid require 3.47 mL KOH for neutralization. The volume (in liters) and the molarity of potassium hydroxide and Equation (8) can be used to determine the number of moles of acid present:

$$0.00347 \text{ L} \times \frac{0.6007 \text{ mol KOH}}{\text{L}} \times \frac{1 \text{ mol HA}}{1 \text{ mol KOH}} = 0.0020\underline{8}1 \text{ mol HA}$$
(9)

Laboratory Technique for Burets

Burets are used to deliver a recorded amount of liquid or solution to another container. A buret is marked in milliliters like a graduated cylinder, but buret markings show **0 mL** at the top, and the numbers increase as you go down the buret. The stopcock controls the liquid flow. It is *open* when *parallel* to the length of the buret and *closed* when *perpendicular* to the length of the buret.

• Washing and rinsing the buret: To clean a buret, wash its interior with soap and tap water. Next, rinse the buret with 5-10 mL portions of DI water. With the buret over the sink and the stopcock open,

• **Conditioning the buret:** After the buret is well-drained, close the stopcock and add about 5 mL of the *titrant* (the solution to be used into the buret). Tilt the buret sideways and roll the barrel to completely rinse the inner walls of the buret. Drain the solution through the buret tip to insure the tip is also conditioned. Repeat this step at least twice to be sure all interior surfaces are rinsed with titrant.

• **Filling the buret:** Close the stopcock. Use a clean funnel to fill the buret with titrant just above the "0" mark. Place a container under the buret tip, and open the stopcock briefly to fill the buret tip with solution, leaving no air bubbles, and to get the level of meniscus to fall within the markings of the buret. If the tip does not fill with solution when the stopcock is in the open position, there may be an air bubble in the stopcock. Consult your instructor.

Note: *The initial level of titrant need not be exactly at 0.00 mL* as the initial level of liquid will be recorded and subtracted from the final volume to determine the volume delivered.

• **Reading the buret:** Always remove the funnel used to fill the buret before taking any measurements. Record the volume of titrant by noting the bottom of the meniscus. On the buret shown below, numbers marked for every 1 mL, and the ten lines between each number represent every 0.1 mL. Thus, the level of titrant in the buret can be estimated to one more decimal place than the markings or to the nearest 0.01 mL.

Thus, in the figure to the right, the meniscus is about halfway between 25.0 and 25.1 mL, so the level of titrant can be recorded as 25.04 mL, 25.05 mL, or 25.06 mL depending on whether the bottom of the meniscus appears to be just above, just at, or just below halfway, respectively.



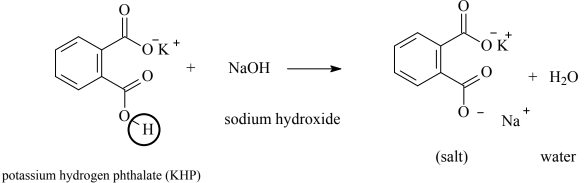
• **Cleaning the buret:** Afterwards, empty the buret, disposing of the titrant according to the waste disposal instructions for each experiments. Wash the buret with soap and tap water, then rinse with several portions of tap water, allowing some tap water to run through the tip. Do a final rinse with small portions of DI water, allowing the DI water to run through the tip, then return the buret to the stockroom.

Part I: Standardization of NaOH with KHP

In this experiment the primary standard used will be **potassium hydrogen phthalate**, **KHC**₈**H**₄**O**₄, which is abbreviated as **KHP**. (Note that the "P" in KHP is *phthalate*, not phosphorus!) KHP is used to standardize the sodium hydroxide solution. The balanced chemical equation for the reaction is shown below:

 $\operatorname{KHC}_{8}\operatorname{H}_{4}\operatorname{O}_{4}(aq) + \operatorname{NaOH}(aq) \rightarrow \operatorname{H}_{2}\operatorname{O}(l) + \operatorname{NaKC}_{8}\operatorname{H}_{4}\operatorname{O}_{4}(aq)$ (3)

(The reacting hydrogen is circled in the equation below.)



otassium hydrogen phthalate (KHP) (KC₆H₈O₄) In this experiment, phenolphthalein will be used as the indicator because it gives a sharp endpoint, meaning that when a titration is being carried out, the phenolphthalein is so sensitive that within a fraction of a drop of titrant, phenolphthalein will make the solution turn completely pink. Note that phenolphthalein does not turn pink until the solution is *basic*, so at the endpoint, when the color change occurs, there is a slight excess of hydroxide ions in solution, making the solution basic. The fainter the pink color, the closer one is to the equivalence point (also called the stoichiometric point), the theoretical point in the titration when enough titrant has been added to react completely with the analyte.

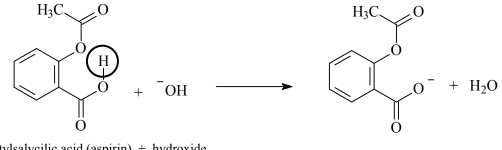
Part II: Determination of the purity of aspirin tablets

Second, you will titrate a sample of your aspirin (acetylsalicylic <u>acid</u>) with the standardized NaOH to determine the moles of acid in a given weight of your product. This will allow you to assess its purity.

 $\frac{\text{actual moles of asprin}}{\text{theoretical moles of asprin}} \times 100\% = \% \text{ purity}$

Assuming the aspirin is not contaminated with other acids, the titration allows you to quantitatively determine the purity of your aspirin. (The reacting hydrogen is circled in the equation below.)

The Net Ionic Equation for the titration in this experiment:



acetylsalycilic acid (aspirin) + hydroxide (C₉H₈O₄) (analyte) (titrant) *** The titrant is standardized in Part I of this experiment***

Safety Precautions



NaOH is corrosive. Handle with care. In case of contact with skin, rinse the area with large amounts of water and notify your instructor. Wear goggles at all times in the chemistry laboratory.

Potassium hydrogen phthalate, KHC₈H₄O₄ (abbreviated as KHP) is

an irritant and slightly corrosive. NEVER touch the sample, and be careful transferring the sample to/from the balance. Always use weighing paper, and never place the sample directly on the balance pan, or it will damage the balance! Any KHP spilled on the lab bench must be cleaned up immediately and disposed of in the sink with plenty of water.

Procedure

Part I : Standardization of the NaOH solution

- 1. Review the Laboratory Techniques section on using a buret. Obtain and prepare a buret for titration by washing and conditioning it with titrant solution (NaOH). Fill the buret, eliminate air bubbles, and have your instructor check for air bubble before starting your first titration. Your buret is now ready for the rest of the lab (**NOTE: You should not need more than 50 mL NaOH for all the titrations**)
- 2. Record the initial buret reading to the correct number of decimal places. *Don't forget to take the funnel out before all readings!*
- 3. Locate a small vial of KHP near the balance area. Tare a folded weigh-paper, and remove it from the balance pan. Use a metal spatula to transfer a small amount of KHP to the paper, and then return the paper to the balance. Remove the paper to add KHP if necessary. *NEVER RETURN CHEMICALS TO THE STOCK CONTAINER!* Record the mass, and transfer the KHP to the flask, rinsing the paper with a gentle stream of DI water to ensure complete transfer.

Place approximately 0.8 g of KHP (MW = 204.23 g/mol) in a clean 125 mL Erlenmeyer flask. Record the precise mass used. Dissolve the acid in 50 mL of distilled water and add 3 drops of phenolphthalein indicator.



Transfer the estimated volume of solution required to react completely with the KHP from Pre-lab question #4 to the box on the right. Use this approximation to determine when your titration will approach the endpoint, and slow addition of NaOH to one drop per second as this volume is reached.

mL (approx. volume required)

- 4. Run slightly less (1-2 mL less) than this amount into the flask containing the acid, while swirling the flask. Use your wash bottle to clean the walls of the flask of drops of base that may have splattered out the titration mixture. Continue adding the NaOH solution dropwise, swirling the flask constantly to ensure full mixing. As you approach the volume estimated for complete reaction, add the NaOH more slowly, (one drop at a time) while continuing to swirl the flask and wash down the walls of the flask.
- 5. Stop the titration when the addition of <u>a single drop</u> of NaOH changes the color of the solution to a light pink, indicating that you have reached the endpoint. The endpoint should persist for 30 seconds without fading.
- 6. Record the final buret reading on you data table.
- 7. **Repeat the titration**. Your molarity should reflect the most significant figures you can obtain with the lab equipment, and the molarity obtained from two runs should agree within 5%. For each run:
 - Calculate the molarity of your NaOH solution.
 - Average the molarity of the first two runs, \overline{M} .
 - Compute the percent difference between the two runs:

 $\circ \quad \frac{|\mathbf{M}_1 - \mathbf{M}_2|}{\overline{M}} \times 100\% = \% \text{ difference}$

- o If run 1 and run 2 are within 5% agreement (of molarity), proceed to Part 2.
- If your runs are not in close agreement, run a third trial. Find the percent difference between the two closest runs, and report this in your Summary Table.
- o If you omit a run explain why (i.e. over titrated sample; passed the endpoint).

Think about it! If your percent difference is within 5% what does that mean about your data?

Part II: Titration of the Aspirin Sample

- 1. Obtain two tablets of commercially available aspirin, weigh them, and place them into a 250-mL flask. Record the brand name to identify your unknown.
- 2. Obtain about 50 mL of 50% ethanol solution and add it to the flask.
- 3. Allow the tablets to stand for a few minutes, and then tap the tablets using a glass stirring rod. They should disintegrate, and the solution will remain cloudy due to the insoluble starch binder. Swirl the mixture to dissolve the aspirin. Aspirin itself does not dissolve well in water-- an ethanol solution is used to dissolve the sample.
- 4. Add several drops of phenolphthalein indicator. Slowly titrate the aspirin with the standardized NaOH solution. Record the initial and final buret readings to the correct number of significant figures. Calculate the percent purity of your sample.
- 5. **Repeat the titration**. Calculate and report the percent purity of your aspirin sample. Do a third trial if percent difference in your percent purity is greater than 5% and time allows. Report the moles of acid found and percent difference in percent purity of the closest two trials.

 $\frac{\text{actual moles of asprin}}{\text{theoretical moles of asprin}} \times 100\% = \% \text{ purity}$

Waste Disposal: All solutions should be emptied into appropriate containers in the fume hood.

Report Sheets Titration of Aspirin Tablets

Name

Lab partner_____

Section____

Background

Calculate the molar mass of KHP and enter it here:

Show the calculation for Pre-lab Exercise #4, estimating the volume of 0.1 M NaOH required to neutralize 0.8 g KHP.

Data

Table 1: Standardization of NaOH

	Trial 1	Trial 2	Trial 3	Trial 4
Mass KHP (g)				
Moles KHP				
Moles NaOH				
Final volume (mL)				
Initial volume (mL)				
Volume of NaOH delivered (mL)				
Molarity of NaOH				
Average Molarity				
% difference				
Observations:				

Don't forget to use units and significant figures.

Report Sheets

Record the brand name of the unknown aspirin sample:

Table 2. Determination of acetylsancy	ylic acid in aspirin samples (purity)			
	Trial 1	Trial 2	Trial 3	
Mass aspirin tablets (g)				
Final volume (mL)				
Initial volume (mL)				
Volume of NaOH delivered (mL)				
Moles of NaOH delivered				
Moles of aspirin				
grams aspirin in tablets				
percent purity of aspirin				
average purity		1	1	
Observations:	a+			

Table 2: Determination of acetylsalicylic acid in aspirin samples (purity)

Don't forget to use units and significant figures.

Calculations

Part I: Standardization of the NaOH solution

- a) Calculate the moles of KHP used in each trial.
- b) Calculate the volume of NaOH required to neutralize the KHP in each trial.
- c) Calculate the molarity of the NaOH for each trial.

Report Sheets

d) Average these values to find the average molarity (M) that will be used in part II.

Average Molarity of standardized NaOH (M):

e) Find the percent difference for the closest two trials, and report this value (also indicate which trials were included) in Table 2. (This should be done in lab, before moving on to Part II.)

Part II: Titration of the Aspirin Sample

- a) Assuming that your aspirin tablets are pure aspirin, calculate the moles of aspirin for each trial. These moles are the **theoretical** moles of aspirin (in theory, your sample should be pure).
- b) Calculate the moles of NaOH used in each trial.
- c) Calculate the moles of aspirin that reacted with the NaOH in each trial. These are the **actual** moles of aspirin in your sample (the moles of acid that actually reacted with the NaOH).

d) Calculate the percent purity of aspirin in each trial, and report the average percent purity.

Report Sheets

Results

Table 3: Summary of Results for NaOH Standardization				
Part I:	Molarity NaOH	Average molarity	Percent difference	
Trial #				
Trial #				

Table 4: Summary of Results for Aspirin Titration

Part II:	mass sample (g)	mL NaOH delivered	moles aspirin	percent purity	percent difference in purity
Trial #					
Trial #					

Summarize your results using full sentences (in the past tense, third person), and restate the values obtained and the percent differences. What are some sources of error inherent to the techniques used? What potential sources of error could affect a future experimenter?

Post-lab Questions:

1. Give one plausible explanation for why a student might achieve less than 100% yield. (Assume calculations are correct.)

2. Give one plausible explanation for a student who obtains over 100% yield. (Assume calculations are correct.)

Pre-Lab Assignment: Purity of aspirin tablets by titration

Name		
Section		

Read the lab instructions several times. Refer to the sections in your textbook on acid-base titrations.

- 1) Sodium hydroxide is a hygroscopic solid. It can be purchased as solid pellets. Why can't you weigh out sodium hydroxide on the balance and make a solution of known concentration? (In other words, why does NaOH need to be standardized?)
- 2) When mixed HCl an acid and NaOH a base will react with each other.
 - a) Write the balanced neutralization equation for this reaction
 - b) What volume of 0.812 M HCl is required to titrate 1.33 g of NaOH to the endpoint?

 A student found that his titration had taken 10.00 mL of 0.1002 M NaOH to titrate 0.132 g of aspirin. Calculate his percent purity. Give a possible explanation of what might have affected his percent purity.

4) The NaOH solution is standardized with KHP in part 1 of this lab. If about 0.8 grams of KHP (MW = 204.22 g/mol) is titrated with approximately 0.1 M NaOH, what is the approximate volume (mL) of NaOH required to reach the endpoint? Show your calculation below. Write this value in the box located in the procedure section for Part 1.