

Presented By : Mustafa Kharma

a BYJU'S

### (تحضير وتحليل الأسبرين) Experiment 7: Aspirin Synthesis and Analysis

#### **Objectives:**

- ► To synthesize aspirin .
- ► To determine the purity of the synthesized aspirin or a commercial aspirin tablet.

Please the video (synthesis of aspirin) at the link

https://youtu.be/oSk-2CQM6zE

Aspirin is a leading commercial pain reliever, first synthesized in a pure and stable form by

Felix Hoffman in 1897.

### ACETYLSALICYLIC ACID STRUCTURE







#### Introduction: Chemical background

Pure aspirin, chemically called acetylsalicylic acid, is both an organic ester and an organic acid. It appears as a white crystalline powder.

► <u>Aspirin</u> is one of the safest and most effective medicines and is widely used medication, thus is displayed on the WHO's List of Essential Medicines.

It is used widely as a <u>painkiller (دواء مسكن للألم</u>) analgesic) such as headache, antipyretic <u>(خافض للحمى</u>) as a fever-reducing drug. It is most widely used in medication to treat pain, inflammation, and fever.

► When ingested, acetylsalicylic acid (ASA) remains intact in the acidic stomach, but in the basic medium of the upper intestinal tract (الجهاز الهضمي العلوي) it forms the salicylate and acetate ions .

► The analgesic action (عمل مسكن ) of aspirin is due to the salicylate ion.





+ H<sub>2</sub>C

Aspirin (acetylsalicylic acid)

Salicylate ion

Acetate ion



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#### Question. Identifthre medical applications of aspirin.

Select one or more:

- a) Fever-reducer
- b) Pain killer
- c) Anti- coagulant
- d) Anti-histamine
- e) Anti -inflammability

#### Determination of the purity of an aspirin sample or commercial aspirin tablet

#### 1) Determination of melting point of ASA.

Qualitative analysis, the purity of an aspirin sample can be determined from its melting point (135°C for pure aspirin). The melting point of a substance is essentially independent of atmospheric pressure, but it is always lowered by the presence of impurities (a colligative property of pure substances. The degree of lowering of the melting point depends on the nature and the concentration of the impurities.

#### 2) Determination by acid-base titration:

Direct titration of ASA with standard solution of NaOH to the endpoint of indicator
 Phenolphthalein indicator : an acid-base indicator that is colorless at a pH less than 8.2 and pink at a pH greater than 10.



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#### **Titration of ASA with NaOH titrant**



#### At the endpoint of titration:

- ► Since, one mole of NaOH reacts with one mole of acetylsalicylic acid (ASA or 1 mol ASA/1 mol OH-), then mol acetylsalicylic acid (ASA) = mol NaOH X  $1/1 = V_{NaOH} \times M_{NaOH}$
- ▶ Mass of ASA (g) = mol ASA x 180.2 g ASA /1 mol ASA ≻
- purity (m/m) = (g ASA/g aspirin sample) x 100 %

#### **Preparation Of Aspirin**

### la Medica

Aspirin (180.2 g/mol, ) is prepared by reacting salicylic acid (138.1 g/mol) with acetic anhydride (102.1 g/mol). Aspirin, like many other organic acids, is a weak monoprotic acid (حمض احادي البرتون).





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### ( الأجراء العملي) Experimental Procedure

#### Procedure overview :

The melting point. Crystalline aspirin is synthesized and then purified by recrystallization and the percent purity of the aspirin are determined, the latter by titration with a standardized NaOH solution.

▶ Be aware of the number of significant figures when recording data.

#### A. Aspirin (Preparation) synthesis

It is safest to prepare the aspirin in a fume hood ( خزانة الأبخرة ).

#### Step 1. Mix the starting materials and heat

▶ weigh ~2 g (±0.01 g) of salicylic acid in a dry 125-mL Erlenmeyer flask. Add 4–5 mL of acetic anhydride. (Caution: Acetic anhydride is a severe eye irritant—avoid skin and eye contact.), then swirl the flask to wet the salicylic acid crystals.

► Add 5 drops of conc H2SO4 (Caution: H2SO4 causes severe skin burns) to the mixture and gently heat the flask in a boiling water bath for 5–10 minutes. H2SO4 is added as a catalyst which speed up the reaction.



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#### Boiling water bath for the dissolution of the acetylsalicylic acid crystals

#### Step 2. Cool to crystallize the aspirin

- ► Remove the flask from the hot water bath, add ~10 mL of deionized ice water to decompose any excess acetic anhydride in the reaction mixture.
- ► Keep the flask in an ice bath to cool the mixture and to speed crystallization. Stirring infrequently to decompose residual acetic anhydride.

#### Step 3. filtration, and washing of solid aspirin

- ► Set up a vacuum filtration apparatus (جهاز ترشيح الفراغ) and turn it on (next Figure). Seal the filter paper with water in the Büchner funnel (قمع بوخنر).
- the mixture onto the filter. Repeat until the transfer of the crystals to (, /سكب/decant) Pour the vacuum filter is complete.
- ► Wash the aspirin crystals on the filter paper with 10 mL of ice-cold water to minimize the loss of the product
- ► Maintain the vacuum for a while to dry the crystals.



#### Step 4. Recrystallize the aspirin: to purify the aspirin crystals

Transfer the crystals from the filter paper(s) to a 100-mL beaker. Add 10 mL ethanol. Warm the mixture in a 60°C water bath (Caution: No flame—use a hot plate or a hot water bath).
Pour 50 mL of 60°C water into the solution and heat the solution to dissolve the solid, do not boil.

Cover the beaker with a watch glass, remove it from the heat, and set it aside to cool slowly to room temperature. Then set the beaker in an ice bath. Beautiful needlelike crystals of acetylsalicylic acid form (تشبه الإبرة بلورات).



#### Step 5. filtration. Vacuum filter the crystals on filter paper

Wash the crystals with two 10-mL volumes of ice water. Place the filter paper and aspirin sample on a watch glass and allow them to air-dry.

#### Step 6. Correct for residual solubility.

The solubility of acetylsalicylic acid is ~0.25 g per 100 mL of water. Correcting for this inherent loss of product due to the wash water. Weigh the aspirin crystals, this is the Experimental yield of aspirin. Calculate the percent yield

#### percent yield = (theoretical yield (g)/ actual yield (g) )x100 %

#### B. Melting Point (m.p) of the Aspirin Sample

The m.p of the aspirin sample can be determined with either a commercial. m.p apparatus or with the apparatus shown in next Figure.

#### Step 1. Prepare the sample.

Fill a capillary m.p tube to a depth of 1 cm with the recrystallized aspirin.

Attach the tube to a 360°C thermometer with a rubber band. As the m.p for aspirin is greater

than 100°C, a cooking oil must be used for the heating bath.



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#### **Step 2.** Determine the melting point.

Gently heat the oil bath at a rate of ~5°C / minute until the aspirin melts. (**Caution**: The oil bath is at a temperature greater than 100°C—do not touch!). Cool the bath and aspirin to just below this approximate m.p until the aspirin in the tube solidifies; at a slower ~1°C /minute rate, heat again until it melts; this is the m.p of your aspirin.



### <u>Step 3.</u> A purity check of the sample (فحص نقاء العينة).

If the m.p of your prepared aspirin sample is less than 130°C, repeat recrystallize the sample for the purpose of increasing its purity.

#### Step 4. Repeat the m.p measurement.

Cool the bath and aspirin to just below the m.p until the aspirin in the tube solidifies; at a 1°C per minute rate; heat again until it melts.



C. Aspirin Analysis: % Acetylsalicylic acid in the Aspirin tablet

#### **<u>Step 1.</u>** Prepare the Aspirin Sample for Analysis.

Crush commercial aspirin tablet (500 mg) and transfer it to clean 250-mL Erlenmeyer flask. Add 10 mL of 95% ethanol, then, 50 mL of deionized water, and swirl to dissolve the aspirin. Add 2 drops of phenolphthalein indicator.

#### Step 2. titrate the sample.

- ► Rinse the clean buret and fill it with a standardized 0.1 M NaOH solution. Record the exact molarity of standard NaOH solution on the report sheet.
- ► Slowly add the NaOH solution from the 50.00 mL buret to the aspirin sample until the endpoint is reached (faint pink color). The color should persist for 30 seconds. Read and record the final volume of NaOH in the buret.

▶ **Repeat the titration**. Three trials are to be completed in the analysis of the aspirin.





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#### **Step 3.** chemical stoichiometry

- ► Calculate the mass of acetylsalicylic acid (g) for the titrimetric analysis.
- ► Calculate the percent purity of aspirin sample (%)
- Calculate the average percent purity of aspirin sample (%)

<u>CLEANUP</u>: Discard the NaOH titrant into a properly labeled bottle; rinse the buret with several 5-mL volumes of tap water, followed by two 5-mL volumes of deionized water.

<u>Caution</u>: 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.





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**Experiment 8. Prelaboratory Assignment. Aspirin Synthesis and Analysis** 

\*\*\* In the experiment, 2.00 g of salicylic acid (molar mass = 138.1 g/ mol) reacts with an excess amount of acetic anhydride.

a. Calculate the theoretical yield of acetylsalicylic acid (molar mass = 180.2 g/mol) for this synthesis.

b. After completing the Experimental Procedure, a mass of 1.78 g of acetylsalicylic acid was recovered. What is the experimental yield for its synthesis?

3. Experimental Procedure, Part B.3. The melting point of the prepared aspirin in this experiment will most likely be less than (but not greater than) that of pure aspirin. Explain.

4. A 0.421-g sample of aspirin prepared in the laboratory is dissolved in 95% ethanol, diluted with water, and titrated to the phenolphthalein endpoint with 17.3 mL of 0.114 M NaOH.

## .a medica

a. How many moles of acetylsalicylic acid (molar mass 180.2 g/mol) are present in the sample?

b. Calculate the percent purity of acetylsalicylic acid in the aspirin sample.

5. 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.