Instruction Hours/week:L: 0 T:0 P:2

Marks: Internal:40 External: 60 Total:100

Course Objectives

It enables the students

- 1. To test the organic functional groups like alcohols, phenols carbonyl and carboxylic acid groups.
- 2. To carryout the preparations of organic compounds by acylation reactions.
- 3. To carryout the preparations of organic compounds by benzyloation reactions.
- 4. To carryout the iodoform reactions and selective reductions.
- 5. To prepare semicarbazone derivatives of ketones.

Course Outcome

The student know to

- 1. Identify the organic functional groups like alcohols, phenols carbonyl and carboxylic acid groups.
- 2. Prepare organic compounds by acylation reactions.
- 3. Prepare organic compounds by benzoylation reactions.
- 2. Carryout the iodoform reactions and selective reductions.
- 3. Prepare semicarbazone derivatives of ketones.

Experiments

- 1. Functional group tests for alcohols, phenols, carbonyl and carboxylic acid group.
- 2. Organic preparations:
- i. Acetylation of one of the following compounds: amines (aniline, o-, m-, p- toluidines and o-, m-, p-anisidine) and phenols (β -naphthol, vanillin, salicylic acid) by any one method:
 - a. Using conventional method.
 - b. Using green approach
- Benzolyation of one of the following amines (aniline, o-, m-, p- toluidines and o-, m-,panisidine) and one of the following phenols (β -naphthol, resorcinol, p- cresol) by Schotten-Baumann reaction.
- iii. Oxidation of ethanol/ isopropanol (Iodoform reaction).
- iv. Selective reduction of meta dinitrobenzene to m-nitroaniline.
- v. Hydrolysis of amides and esters.
- vi. Semicarbazone of any one of the following compounds: acetone, ethyl methyl ketone, cyclohexanone, benzaldehyde.

- vii. Esterification of any one of the following acids (benzoic acid, oxalic acid, phenyl acetic acid and phthalic acid).
- viii. Aldol condensation using either conventional or green method.

The above derivatives should be prepared using 0.5-1g of the organic compound. The solid samples must be collected and may be used for recrystallization and melting point.

Suggested Readings

Text Books:

- 1. Mann, F.G. & Saunders, B.C. (2009). *Practical Organic Chemistry*. Pearson Education.
- 2. Furniss, B.S., Hannaford, A.J., Smith, P.W.G. & Tatchell, A.R. (2012). *Practical Organic Chemistry* (V Edition). Pearson.
- 3. Venkateswaran, R., Veeraswamy, R. & Kulandaivelu, A.R. (2015). *Basic Principles of Practical Chemistry*. S. Chand & Sons Ltd., New Delhi.
- 4. Bansal, R.K. (2012). *Laboratory Manual of Organic Chemistry* (V Edition). New Age International Publishers (P) Ltd, New Delhi.
- 5. ThomasA.O. (2003). *Practical Chemistry for B.Sc Main Students*, Scientific Book Centre, Cannore-1, Kerala.

Reference Books:

1. Ahluwalia, V.K., & Aggarwal, R. (2000). *Comprehensive Practical Organic Chemistry: Preparation and Quantitative Analysis*. University Press.

COURSE CODE: 18CHU213

BATCH-2018-2021

PREPARATIONS

1. ACETYLATION OF ANILINE TO ACETANILIDE

PRINCIPLE:

Aniline can be acetylated easily by reflecting a mixture of acidic anhydride and sodium acetate

 $CH_6H_5NH_2 + (CH_3CH_2)_2 O + CH_3COONa \rightarrow C_6 H_5 NH COCH_3 + CH_3 COOH$

REQUREMENTS:

Chemicals:

 $C_6H_5NH_2$ (freshly prepared) -3ml

Acetic anhydride -- 4ml

Glacial acetic acid --4ml

APPARATUS REQUIRED:

Reflex conteser, glass beaker (250ml), wire gase, test tube (10ml) and round bottle flask

PROCEDURE:

Galacial acetic acid is mixed with acidic anhydroxide in dry and tested an shaken well and aniline is taken in round bottom flask and the above mixture is added slow to aniline with contantshaking. A bit of broken borceline is added into the flask to prevent pumping. The flask is fitted with a reflex condenser and heated gently for above 30min over a wire gasze the hot liquid is poured into ice cold water taken in formed product is filtered, dryed and the yield is noted.

RECYSTILLATION:

About 1g of the crud acetanilide sample is dissolve in minimum amount of hot water, 1g of animal char coal is added and then heated the hot liquid is rapidly

KARPAGAM ACADEMY OF HIGHER EDUCATIONCLASS: I B SC CHEMISTRYCOURSE NAME: OXYGEN CONTAININGFUNCTIONAL GROUP-PRACTICALEATCH-2018-2021COURSE CODE: 18CHU213BATCH-2018-2021

filtered (through hot water, fume is available). The filtered allowed to cool, the acetanilide crystallize in a rhompic plates.

RESULT: 1. MELTING POINT: 2. EXPECTED YIELD:

KARPAGAM ACADEMY OF HIGHER EDUCATIONCLASS: I B SC CHEMISTRYCOURSE NAME: OXYGEN CONTAININGFUNCTIONAL GROUP-PRACTICALEATCH-2018-2021COURSE CODE: 18CHU213BATCH-2018-2021

2. PREPARATION OF BENZAMIDE FROM ANILINE

AIM

To prepare a few sample of crystals of Benzamide from aniline.

PRINCIPAL

Aromatic amines react with benzyl chloride in aqueous medium to yield the respective benzanilides. In the Scotten Burman method of benzoylation's amino compound is treated with a site excess of NaOH solution and benzyl chloride, on vigorous shaking. Benzylation taken place radically and the product separates as a solid.

 $C_6H_5NH_2 + C_6H_5COCl + NaOH - C_6H_5NHCOC_6H_5 + NaCl + H_2O$

CHEMICAL REQUIRED

Aniline - 2.5 ML

Benzyl chloride - 3.5 ML

NaOH - 10% 25ML

PROCEDURE

2.5ML of aniline and 25ml of 10% NaOH are placed in a 100ml conical flask and stopped well. 3.5ml of benzyl chloride are introduced into to the flask vigorously shaken well for 10-15 mints. The progress of reaction is known by the exothermal process. The completion of reaction is tested by smelling the presence of benzyl chloride in the flask. If the smell precisive add little more NaOH and shake for a while. The white crystals of benzyl chloride are filtered at the pumb washed well with H2O and dried.

RECRYSTALLISATION

A little amount of the sample is recrystalized from hot methylated split it. It is filtered to hot water funnel. The colourless crystals are separated and dried in air.

RESULT

The yield of benzylanilide -

3. Preparation of benzoic acid from ethyl benzoate

Principle:

Ethyl benzoate is hydrolysed to sodium salt of benzoic acid by a solution of sodium hydroxide. Benzoic acid is obtained from sodium benzoate by acidification.

Step:1

 $C_6H_5COOC_2H_5 + NaOH \rightarrow C_6H_5COONa$

Step:2

 $C_6H_5COONa + HCl \rightarrow C_6H_5COOH$

Requirements:

- 1. Ethyl benzoate-2.5g
- 2. Sodium hydroxide-2g

Procedure:

2g of sodium hydroxide are dissolved in about 20 ml of water taken in a R.B flask. 2.5 g of ethyl benzoate are added to the R.B flask. A few porous pieces are added to ensure smooth boiling. The flask is then fitted with a Liebig's condenser and heated over a wire gauze for about 45 mins. The hydrolysis is complete when no more oily drops are seen in the flask. The contents are now cooled and transferred completely to a beaker. Concentrated HCl is added with constant stirring till the solution is distinctly acetic. The precipitated benzoic acid is filtered, washed and dried. The yield is noted. About 1g is recrystallized from hot water and melting point is determined.

Result:

- 1. The yield of benzoic acid =..... g
- 2. The melting point of benzoic acid =....°C

4. Oxidation of Iodoform

Principle:

Iodoform is prepared by the joint action of iodine and alkali upon ethanol.

 $\mathrm{CH_3CH_2OH} + 3\mathrm{I_2} \rightarrow \mathrm{CI_3\text{-}CH_2OH} + 3\mathrm{HI}$

 CI_3 - $CH_2OH + NaOH \rightarrow CHI_3 + CH_3COONa$

Requirements:

Ethanol- 3.5ml

Potassium carbonate- 5g

Powdered iodine- 5g

Procedure:

The Potassium carbonate is dissolved in 20ml of water contained in a 100ml conical flask. Ethanol is added to the solution. The powdered iodine is then added in small quantities at a time with stirring.

The mixture is gradually warmed on a water bath to $70 - 80^{\circ}$ C for about 15 mins. On cooling, the iodoform seperates as yellow crystals and is collected by filteration. The product is washed with water, recrystallized from alcohol and dried. The yield and melting point is determined.

Result:

- 1. The yield of iodoform =..... g
- 2. The melting point of iodoform = $\dots^{\circ}C$

5. Preparation of m-Dinitrobenzene

Principle:

m-Dinitrobenzene is prepared by the nitration of nitrobenzene. The nitration is effected by fuming nitric acid in presence of conc.H2SO4.

Requirements:

nitrobenzene- 1ml

Fuming nitric acid- 5ml

Conc. H₂SO₄-10ml

Procedure:

7ml of fuming nitric acid is taken in an R.B flask and 10ml of Conc. H_2SO_4 is added, a little at a time, cooling the flask during the addition. 1ml of nitrobenzene is then added in small quantities at a time to the nitrating mixture, shaking well after each addition. Finally it is heated for about 45mins by immersion in a boiling water-bath, till a small quantity of the reaction mixture when added to small quantity of water in a test-tube gives a solid immediately. The contents of the flask are then poured in fine stream while still hot into 100ml of water contained in a beaker. The mixture is shaken very vigorously during the addition. M-Dinitrobenzene separates as a solid which is filtered at the pump, washed several times with water, dried and the yield noted. A small portion is recrystallized from alcohol and the melting point is determined.

Result:

- 1. The yield of m-dinitrobenzene =..... g
- 2. The melting point of m-dinitrobenzene =....°C

CLASS: I B SC CHEMISTRY COURSE NAME: OXYGEN CONTAINING FUNCTIONAL GROUP-PRACTICAL

COURSE CODE: 18CHU213

BATCH-2018-2021

ORGANIC QUALITATIVE ANALYSIS

TESTS FOR FUNCTIONAL GROUPS

١. Compounds in which carbon, Hydrogen and Oxygen are present

I.1. **Carboxylic Acids**

(a)	A pinch of the substance is	A pink colour is produced	Presence of carboxylic
	shaken up with about 1 ml of	only after adding a number	acids.
	water. One drop of	of drops of sodium	
	phenolphthalein is added and	hydroxide.	
	then very dilute sodium		
	hydroxide solution is added	A pink colour is got even	Acids absent
	drop by drop with shaking.	with the first drop of sodium	
		hvdroxide.	
		3	
(b)	Ester formation:	A pleasant ester smell is	Carboxylic acid is
	About 0.2 gm of the	obtained.	present.
	substance is mixed with		
	about 2 ml of rectified spirit		
	in a dry test tube: 2 to 3 drops		
	of conc. H_2SO_4 are added.		
	shaken well and gently		
	heated for a minute The		
	mixture is then poured into		
	about 30 ml of a dilute		
	solution of sodium carbonate		
	stirred well and the smell is		
	noted		
	hours.		
(c)	Fluorescein reaction:	The mixture turns deep-red	Dicarboxylic acids
	A small amount of the	on heating; forms a red	present.
	substance is mixed with a	solution with water: on	1
	few crystals of resorcinol in a	adding sodium hydroxide, an	
	dry test tube. 2 drops of conc.	intense greenish yellow	
	H_2SO_4 are added, shaken	fluorescein is obtained.	
	well, heated gently and then		
	poured into about 100 ml of		
	water, stirred well and then		
	an excess of sodium		
	hydroxide solution is added.		

I.2. Phenolic Group

(a) *Libermann's Reaction:*

COURSE NAME: OXYGEN CONTAINING CLASS: I B SC CHEMISTRY FUNCTIONAL GROUP-PRACTICAL COURSE CODE: 18CHU213

BATCH-2018-2021

	A small amount of the	A red solution is got which	Phenolic group present.
	substance is heated with a	turns blue or green on	
	crystal of sodium nitrite in a	adding sodium hydroxide.	X 11 1 1 1
	dry test tube, cooled, two		Note: all phenols do not
	arops of conc. H_2SO_4 are		give this test.
	mixture is poured into about		
	100 ml of water stirred well		
	and then an excess of sodium		
	hydroxide is added.		
	2		
(b)	Phthalien fusion Test :	A pink or red colour is	Phenols like phenol, o-
	A small amount of the	obtained.	cresol, m-cresol,
	substance is mixed with about		salicylic acid.
	0.5 gm of phthalic anhydride		Deservine1
	in a dry tube; 2 drops of	A yellowish green	Resorcinoi
	gently for a minute. The	nuorescence is obtained.	Note: The colour
	mixture is poured into about		depends on the nature of
	100 ml of water and the		the phenol.
	sodium hydroxide is added in		1
	slight excess.		
(c)	Azo-Dye Formation :	A scarlet red or brownish	Phenols like β -naphthol,
	About 5 drops of aniline are	red or orange red precipitate	α -naphthol, resorcinol
	treated with 3 to 5 ml of dil.	of a dye is obtained.	etc.
	solution of sodium nitrite are		
	added drop by drop with		
	constant shaking and cooling.		
	and the diazotised solution		
-	obtained is added to a solution		
	of the substance in 2 to 3 ml of		
	10% NaOH solution in small		
	quantities.		
(d)	About 1 ml of Febling's	A red brown precipitate of	Polyhydric Phenol
(4)	solution 'A' is mixed with 1	cuprous oxide is obtained	i orynyanie i nenoi.
	ml of Fehling's solution 'B'.		
	The mixture is treated with a		
	small amount of the substace		
	and boiled well.		

I.3. Aldehydes and Ketones

(a)	About 3 drops of the substance	A white crystalline	Aldehydes and certain
	are added to a saturated	precipitate is obtained.	ketones present.

Prepared by Dr. K. Sundaram, Assistant Professor, Department of Chemistry, KAHE Page 8/11

CLASS: I B SC CHEMISTRYCOURSE NAME: OXYGEN CONTAININGFUNCTIONAL GROUP-PRACTICALEATCH-2018-2021COURSE CODE: 18CHU213BATCH-2018-2021

	solution of sodium bisulphite solution and shaken well.		[ketones with the keto group directly attached to benzene ring does not answer this test]
(b)	A small amount of the substance is added to about 3 ml of Borsche's reagent, a drop of conc.HCl added, gently heated for about 2 minutes and cooled well.	A yellow or red-brown crystalline precipitate is obtained.	Aldehydes and ketones present.
(c)	A mixture of 5 drops of Phenylhydrazine and 5 drops of glacial acetic acid are taken in dry test tube. A small amount of the substance is added, gently heated for a minute. Excess of cold water is added and shaken well.	 (a) A yellow or yellowish-white precipitate is obtained. (b) A bright yellow crystalline precipitate. 	Presence of Aldehydes and certain Ketones. Reducing sugars present.
(d)	About 0.3 gm of semicarbazide hydrochloride id dissolved in H_2O . About 0.5 gm of sodiumacetate crystals are added, shaken well to dissolve the solid and to this 2 ml of an alcoholic solution of the substance is added, heated in a water bath for 15 minutes and cooled.	A white crystalline precipitate of the semicarbazone is got.	Aldehydes and Ketones present.
(e)	A small amount of the substance is added to about 3ml of Schiff's reagent and shaken well.	A violet colour is produced quickly.	Aldehydes confirmed.
(f)	A small amount of the substance is added to 2 ml of Fehling's solution A and B and gently heated.	A reddish brown precipitate is got.	Presence of Aliphatic aldehydes (reducing sugars, and polyhydric phenols also answer).

1.4. Carbohydrates

(a)	Molisch test:	

Prepared by Dr. K. Sundaram, Assistant Professor, Department of Chemistry, KAHE Page 9/11

COURSE NAME: OXYGEN CONTAINING CLASS: I B SC CHEMISTRY FUNCTIONAL GROUP-PRACTICAL COURSE CODE: 18CHU213

BATCH-2018-2021

	A		
	A small amount of the	A violet, purple or red ring	Carbonydrates
	substance is dissolved in 2 ml	is formed at the junction of	confirmed.
	of water, a few drops of a	the two layers and the	
	strong solution of (10%) of α -	colour slowly spreads	
	naphthol in pure alcohol is	throughout the liquid.	
	added, shaken well, and 2 ml		
	of conc. H_2SO_4 are carefully		
	added along the side of the		
	tube so that the acid forms a		
	separate layer at the bottom.		
	- F		
(b)	A small amount of the	A red brown precipitate is	Reducing sugar present
(-)	substance is shaken up with	obtained	
	one ml of water. The solution		
	is added to about 2 ml of		
	Fehling's A and B and heated		
	on a water bath for 15 minutes		
	on a water bath for 15 minutes.		
(c)	About 0.3 gm of the substance	A bright yellow crystalline	Reducing sugars like
(0)	is dissolved in about 5 ml of	precipitate is obtained	glucose fructose and
	water a mixture of 2 ml of	precipitate is obtained.	lactose present
	water, a mixture of 2 mi of		lactose present.
	phenyinyulazine and 2 mi ol		
	glacial acetic acid is added, the		
	contents heated in a boiling		
	water bath for about 15		
	minutes with occasional		
	shaking and then cooled.		

Esters:

(a)	Hydroxamic acid Test:		
	About 3 drops of the liquid and a pinch	A violet or deep red	Esters present.
	of hydroxylamine hydrochloride are	brown colour	
	added to about 5 ml of 10% NaOH	obtained.	
	solution. The contents are gently boiled		
	for 3 minutes cooled, acidified with		
	conc.HCl, added drop by drop and then		
	about 5 drops of FeCl ₃ solution are		
	added and shaken well.		
(b)	The liquid is refluxed with 10% NaOH	A white precipitate is	Ester of an aromatic
	solution. The residue is acidified with	obtained.	acid is present.
	con.HCl and cooled.		

1.5.