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18CHEL16 : ENGINEERING CHEMISTRY LAB

A. LABORATORY INFORMATION

1. Lab Overview

Degree:	B.E	Program:	BS
Year / Semester :	2019/1	Academic Year:	2019-20
Course Title:	Engineering Chemistry Lab	Course Code:	18CHEL16
Credit / L-T-P:	1/0-0-2	SEE Duration:	180 Minutes
Total Contact Hours:	42 Hrs	SEE Marks:	60 Marks
CIA Marks:	40	Test	2
Course Plan Author:	Dr. Manju M	Sign	Dt : 04-01-2019
Checked By:	Dr. Shankara B.S	Sign	Dt : 14-08-2019

2. Lab Content

Unit	Title of the Experiments	Lab Hours	Concept	Blooms Level
	PART- A			
1	Potentiometric estimation of FAS using standard K 2 Cr 2 O 7 solution.		Redox Reaction s	L4 Analyzing & L5 Evaluation
2	Conductometric estimation of acid mixture.		Acid Base Reaction	L4 Analyzing & L5 Evaluation
3	Determination of Viscosity co-efficient of the given liquid using Ostwald's viscometer.		Cohesive Force	L4 Analyzing & L5 Evaluation
4	Colorimetric estimation of Copper.	2	Measurem ent of Optical Density	L4 Analyzing & L5 Evaluation
5	Determination of pKa of the given weak acid using pH meter.	2	PH measure ment	L4 Analyzing & L5 Evaluation
6	Flame photometric estimation of sodium and potassium.	2	Atomizati on	L4 Analyzing & L5 Evaluation
	PART- B			
1	Estimation of Total hardness of water by EDTA method.		Complexo metric titration	L4 Analyzing &

Alwanton S	

A REAL PROPERTY OF	SKIT	Teaching Process	Rev No.: 1.0
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				L5 Evaluation
2	Estimation of CaO in cement solution by rapid EDTA method.	2	Complexo metric titration	L4 Analyzing & L5 Evaluation
3	Determination of percentage of Copper in brass using standard sodium thiosulphate solution.	2	lodometri c titration	L4 Analyzing & L5 Evaluation
4	Determination of COD of waste water.	2	Redox titration	L4 Analyzing & L5 Evaluation
5	Estimation of Iron in haematite ore solution using standard K 2 Cr 2 O 7 solution by external indicator method.	2	Redox titration	L4 Analyzing & L5 Evaluatio
6	Estimation of percentage of available chlorine in the given sample of bleaching powder	2	lodometri c titration	L4 Analyzing & L5 Evaluation

3. Lab Material

Unit	Details	Available
1	Text books	
i	Textbook of Engineering Chemistry with Lab Manual 9th Edition (English, Paperback, Shashi Chawla)	In Lib
ii	Vogel's Textbook of Practical Organic Chemistry (5th Edition) 5th Edition by <u>A.I.</u> <u>Vogel</u> (Author), <u>A.R. Tatchell</u> (Author), <u>B.S. Furnis</u> (Author), <u>A.J. Hannaford</u> (Author), <u>P.W.G. Smith</u> (Author)	In Lib
2	Reference books	
i	G.H.Jeffery, J.Bassett, J.Mendham, R.C.Denney, "Vogel's Tex book of quantitative Chemical Analysis Fifth Edition(New) ,	In Lib
ii	O.P.Vermani & Narula, "Theory and Practice in Applied Chemistry", New Age International Publisers.	In Lib
iii	Gary D. Christian, "Analytical chemistry ", 6 th Edition, Wiley India.	In Lib
ii	Engineering Chemistry Lab manual	In dept
3	Others (Web, Video, Simulation, Notes etc.)	
i	https://sites.google.com/chemistry-laboratory-w.	Available on web
ii	https://science.nrao.edu > Facilities > CDL	Available on web
iii	https://www.acs.org//chemistryclubs//simulati	Available on web
iv	https://www.augusta.edu//chemistryandphysics/	Available on web
V	www.ncl-india.org/	Available on web

4. Lab Prerequisites:

-	-	Base Course:		-	-
SNo	Course	Course Name	Topic / Description	Sem	Remarks
	Code				
1			Titrations/students have done these kind	1	
		Chemistry Lab	of experiments in lower standards.		

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		Instrumental analysis/students have	1	
		studied in theory part regarding these		

experiments. Note: If prerequisites are not taught earlier, GAP in curriculum needs to be addressed. Include in Remarks and implement in B.5.

5. General Instructions

SNo	Instructions	Remarks
1	Never work in the laboratory unless a demonstrator or teaching assistant is	
	present.	
2	Do not throw waste such as match stems filter papers etc. into the sink. They	
	must be thrown into the waste jars.	
3	Keep the water and gas taps closed expect when these utilities are needed.	
4	Never taste any chemical unless instructed to do so and don't allow	
	chemicals to come in contact with your skin.	
5	While working with gases, conduct the experiment in a fume hood.	
6	Keep all the doors and windows open while working in the laboratory.	
	You should know about the hazards and properties of every chemical which	
	you are going to use for the experiment. Many chemicals encountered in	
	analysis are poisonous and must be carefully handled.	
	Sulphuric acid must be diluted only when it is cold .This should be done by	
	adding it slowly to cold water with stirring ,and not vice versa.	
	Reagent bottles must never be allowed to accumulate on the work bench.	
-	They should be placed back in the shelves as and when used.	
10	Containers in which reaction to be performed a little later should be labeled.	
	Working	
	space should be cleaned immediately.	

6. Lab Specific Instructions

SNo	Specific Instructions	Remarks
	Chemical Splash Goggles:	
1	Purchase a pair of chemical safety goggles).	
2	Bring your goggles with you for all laboratory sessions of your chemistry class. You will not be allowed to work in the lab without your goggles	
3	Wear your goggles when anyone in the lab is conducting an experiment.	
	Laboratory Coats:	
4	Purchase a lab coat that fits you well. Lab coats that are too tight or too loose are not safe. Sleeves that are too long should be rolled up.	
5	If your lab coat has not been contaminated with a hazardous substance, you may wash it as you do your other clothing.	
6	If your lab coat becomes contaminated with a hazardous substance, as with any other lab spill, notify your instructor immediately.	
7	Contaminated lab coats will be handled by your instructor as they deem appropriate.	
	Nitrile Gloves:	
8	Nitrile gloves are to be worn only during portions of experiments where specified by the experimental procedure, when instructed by the instructor or supervisor, or when working with substances for which the protocol requires the use of gloves.	
9	Note that nitrile gloves are flammable and will stick to your skin if they burn. Do not wear gloves while working with Bunsen burners.	
10	Do not wear gloves outside the lab. When a chemical comes in contact with a glove, remove the glove immediately and place it in the glove waste.	
11	Do not touch surfaces such as door knobs, computer keyboards, and chairs while wearing Pag gloves.	

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12	Gloves			
	proper			
	<u> </u>	<u> </u>		

13 Dispose of gloves at the end of each experiment in the glove waste containers provided in each lab.

B. OBE PARAMETERS

1. Lab / Course Outcomes

#	COs	Teach.	Concept	Instr	Assessment	Blooms'
		Hours		Method	Method	Level
	PAR	Г- А				
	Handling different types of instruments for quantitative analysis of samples.	21	Instrumental method of analysis	Demons trate	Test	L3
	PAR	Г- В				
	Volumetric analysis of various samples quantitatively.	21	Volumetric analysis	Demons trate	Test	L3
-	Total	42	-	-	-	-

Note: Identify a max of 2 Concepts per unit. Write 1 CO per concept.

2. Lab Applications

SNo	Application Area	CO	Level			
PART- A						
1	1 Potentiometric estimation of FAS using standard K 2 Cr 2 O 7 solution.					
2	Conductometric estimation of acid mixture.	CO1	L3 & L4			
	Determination of percentage of Copper in brass using standard sodium thiosulphate solution.	CO2	L3 & L4			
4	Determination of COD of waste water.	CO2	L3 & L4			

Note: Write 1 or 2 applications per CO.

3. Mapping And Justification

4. Articulation Matrix

(CO - PO MAPPING)

-	Course Outcomes	Program Outcomes												
#	COs	PO1	PO2	PO3	PO4	PO5	P06	PO7	PO8	PO9	PO10	PO11	PO12	Level
	Estimate amount of FAS potentic metrically through redox titrations.		×	×										
_	Calculate amount of acid mixture conducto metrically through neutralization titration.	1	×	×										
	Compute amount of copper bu measuring absorbence using optical method		×	X										
	Determine Pka Value of giver sample using Ph meter.	X	x	x										
	Estimation of co-efficient of viscosity of given organic liquic using ostwald's method.		X	X										

	ning Proce	ess		Rev No	.: 1.0		ĺ
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18CHE27.6 Estimate amount of given × ×	x x						
sample using flame photo metric							
method.							
18CHE27.7 Estimation of hardness of given × ×	x x						
sample by using							
complexometric titrations.							
18CHE27.8 Caluculate the % of CaO in given × ×	x x						
sample by rapid EDTA method.							
18CHE27.9 Estimate the % of copper in given × ×	x x						L3
brass sample by iodometric							ĺ
titration.							ĺ

5. Curricular Gap and Content	

18CHE27.10 Calculate the % of iron in given ×

indicator method.

by redox titrations.

18CHE2711. Estimate

1

ore solution using external

total

impurities of given waste water

U	1				
SNo	Gap Topic	Actions Planned	Schedule Planned	Resources Person	PO Mapping
1					
2					
3					
4					
5					
N.L. I					

х х

Х

х

oxidisable ×

Estimate the % of chlorine in given bleaching powder sample by iodometric method

L3

Note: Write Gap topics from A.4 and add others also.

6. Content Beyond Syllabus

SNo	Gap Topic	Actions Planned	Schedule Planned	Resources Person	PO Mapping
1					
2					
3					
4					
5					
6					
7					
8					
9					
10					
11					
12					
13					
14					
15					

Note: Anything not covered above is included here.

C. COURSE ASSESSMENT

1. Course Coverage

1.000.0000	age				
Unit	Title	Teachi	No. of question in Exam	CO	Levels
BSH Prepared by		Checked by		Approved	



Title: Engineering Chemistry Lab

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		ng Hours	CIA-1	CIA-2	CIA-3	Asg-1	.Asg-2	Asg-3	SEE		
		TIOUIS	PART	- A							
1	Potentiometric estimation of FAS using standard K 2 Cr 2 O 7 solution.	02	1	-	-	-	-	-	1	CO1	L3 & L4
2	Conductometric estimation of acid mixture.	02	1	-	-	-	-	-	1	CO1	L3 & L4
3	Determination of Viscosity co- efficient of the given liquid using Ostwald's viscometer.	02	1	-	-	-	-	-	1	CO1	L3 & L4
4	Colorimetric estimation of Copper.	02	1	-	-	-	-	-	1	CO1	L3 & L4
5	Determination of pKa of the given weak acid using pH meter.	02	1	-	-	-	-	-	1	CO1	L3 & L4
6	Flame photometric estimation of sodium and potassium.	02	1	-	-	-	-	-	1	CO1	L3 & L4
			PART	- B							
1	Estimation of Total hardness of water by EDTA method.	02	-	1	-	-	-	-	1	CO2	L3 & L4
2	Estimation of CaO in cement solution by rapid EDTA method.	02	-	1	-	-	-	-	1	CO2	L3 & L4
3	Determination of percentage of Copper in brass using standard sodium thiosulphate solution.		-	1	-	-	-	-	1	CO2	L3 & L4
4	Determination of COD of waste water.		-	1	-	-	-	-	1	CO2	L3 & L4
5	Estimation of Iron in haematite ore solution using standard K 2 Cr 2 O 7 solution by external indicator method.	02	-	1	-	-	-	-	1	CO2	L3 & L4
6	Estimation of percentage of available chlorine in the given sample of bleaching powder		-	1	-	-	-	-	1	CO2	L3 & L4
-	Total	42	7	8	5	5	5	5	20	-	-

Note: Write CO based on the theory course.

2. Continuous Internal Assessment (CIA)

Evaluation	Weightage in Marks	СО	Levels
CIA Exam – 1	10	CO1,	L3 & L4
CIA Exam – 2	10	CO2,	L3 & L4
CIA Exam – 3	10	CO1 & CO2,	L3 & L4
Other Activities – define – Slip test			L2, L3, L4

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Final C	IA Marks	10	-	-			

SNo	Description	Marks
1	Observation and Weekly Laboratory Activities	05 Marks
2	Record Writing	10 Marks for each Expt
3	Internal Exam Assessment	15 Marks
4	Internal Assessment	40 Marks
5	SEE	60 Marks
-	Total	100 Marks

<u>PART - A</u>

D. EXPERIMENTS

Experiment 01 : Potentiometric estimation of FAS using standard K2Cr2O7 solution.

-	Experiment No.:	1 Mar	ks	Date Planned		Date Conducted	
1	Title	Potention	netric estimatio	n of FAS using	g standard	K 2 Cr 2 O 7 s	olution.
2	Course Outcomes		n of amount of F				
3	Aim		netric estimatio		g standard	K 2 Cr 2 O 7 s	olution.
4	Material/ Equipment Required	 Cal 10r 10c Gla 	gital Potentio met lomel & Pt-electr nl Burette Oml beaker ass rod.	odes			
5		a, PRINCIPLE	Redox titratio	ns can be c	arried out	potentiomet	rically using
	Principle, Concept	platinum-c	alomel electrode	e combination	. For the read	ction:	
			Reduced forr	$n \rightarrow Oxidized$	form + ne ⁻ ,		
		The p	otential, E,	is giv	en by	Nernst	equation,
				0	,		
		$E = E^{o} + \frac{1}{2}$	$\frac{0.0591}{n}\log\frac{[\text{Oxio}]}{[\text{Red}]}$	lized form] uced form]			
		Where, E°	is the standard p	otential of the	e system, an	d [X] represe	nt the molar
		concentrat	ion x.				
		Suppose t	hat, in beaker w	e have acidifi	ed Fe²⁺ solu	ution, and we	add slowly
		K ₂ Cr ₂ O ₇ from	m a burette, then	following rea	ction takes p	olace.	
			6 Fe ²⁺ + Cr ₂ O ₇ ²⁻ -	• 6 Fe ³⁺ + 2Cr ³⁺			
		Before the	equivalence po	oint, the poter	ntial is dete	rmined by th	ne Fe²⁺∕ Fe³⁺
		system.					
		$E = E^o + \frac{C}{2}$	$\frac{0.0591}{n}\log\frac{\left[\mathrm{Fe}^{3+}\right]}{\left[\mathrm{Fe}^{2+}\right]}$	=0.75V+0.0	$1591 \log \frac{[\text{Fe}^3]}{[\text{Fe}^2]}$	+] +]	
		The potent	tial of the solutio	n will be arou	nd 0.75V (sir	nce the contr	ibution from
		the second	d term is negligib	le).			
		After the e	equivalence poir	it, the potent	ial is detern	nined by the	e Cr ₂ O ₇ ²⁻ /Cr ³⁺
		system.					

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	· · ·	$E = E^{o} + \frac{0.0591}{6}$	$-\log \frac{\left[\mathrm{Cr}_2 O_{7^{2-}}\right]}{\left[\mathrm{Cr}^{3^+}\right]}$	$\frac{1}{2} = 1.33V + 0$	$0.00985 \log \left[- \right]$	$\frac{\operatorname{Cr}_2 O_{7^{2-}}}{[\operatorname{Cr}_{3+}]}$				
		The potential of t	he potential of the solution will be around 1.3V							
		At the equivalen	t the equivalence point, the potential is average potential of both systems.							
		Thus, there is an	abrupt increa	ase in poter	itial of the so	lution near the end point.				
6	Procedure,					S) solution into a 100 ml				
		beaker. Add two	test tubes fu	ll of dilute s	ulphuric acic	l. Immerse the platinum -				
		calomel electroc	les assembly	in the solu	tion. Measur	e the potential by adding				
		K ₂ Cr ₂ O ₇ solution	from the bur	ette in incre	ements of 0.g	5 cm ³ . Stir the mixture by				
		blowing the air fo	or 10 seconds	. Measure t	he potential	of the each addition.				
		the equivalence	e point from rmality of the	the grap	h. Knowing	shown in the figure. Find the equivalence point, rmine the amount of FAS				
7	Reaction Equation	6 Fe ²⁺ + C	Cr₂O ₇ ²- → 6 Fe ³	'⁺ + 2Cr ^{3⁺}						
				201						
		Vol.of K₂Cr₂O7 in cm³	E in mv	ΔV	ΔΕ	ΔΕ/ΔV				
		0.0								
		0.5								
8		1.0								
	Observation Table,	1.5								
		2.0								
		2.5								
		3.0								
		3.5								
		4.0								
		4.5								
		5.0								
9	Sample Calculations	$\frac{S}{S}$ $\frac{Calculations:}{(N_1 V_1) K_2 Cr_2 O_7 = .(N_2 V_2)_{FAS}}$ Where V ₁ = Vol. of K ₂ Cr ₂ O ₇ at the equivalence point (from the graph) N ₁ = Normality of K ₂ Cr ₂ O ₇ solution =N (to be given) V ₂ = Vol. of FAS solution = 25 cm ³ N ₂ = Normality of FAS solution =N $\therefore N_2 = \frac{V_1 X N_1}{V_2} = \frac{X}{25} =N$ W k t Mass per dm ³ = Normality x Gram equivalent mass Mass of FAS per dm ³ = Normality of FAS x Gram equivalent mass of FAS $=X 392 g =g$								

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	ght ©2017. CAAS. All rights reserved. Graphs, Outputs	$\frac{\Delta E}{\Delta V}$ Equivalence point (V)			
11	Results & Analysis	Volume of K2Cr2O7 in cm3			
		Mass of FAS present in one dm ³ of solution =g			
12		Application of potentiometry to characterize acid and basic sites in humid substances Testing			
		The Techniques to study complexation reactions at the mineral/water.			
		Interface No indicator is used; instead the potential is measured across the analyte, typically an electrolyte solution.			
13	Remarks				
-	Faculty Signature with Date				

Experiment 02 : Conductometric estimation of acid mixture

-	Experiment No.:	2	Marks		Date Planned		Date Conducted	
1	Title	Condu	actometric e	estimation of	acid mixture			
2	Course Outcomes	Calcul reactio	alculate amount of acid mixture conductometrically through neutralization action					
3	Aim	Condu	ictometric e	estimation of	acid mixture	by using sta	ndard NaOH(1N).
4	Material / Equipment Required		Digital Co Conductiv 10ml Bure 100ml bea Acid mixtu 1N NaOH	ette aker ure	r			
5						t sharp and conductivity ations is the conductivity ainst sodium zation of the		
			NaOH	+ HCl		• • • •	JaCl + H₂O	
			NaOH	+ CH ₃ COOH		CH	H₃COONa + H	2O
The addition of sodium hydroxide to hydrochloric a conductance of the latter because highly mobile H ⁺ ions are mobile Na ⁺ ion. This trend continues till all the H ⁺ ions of HC continuing the addition of NaOH, conductance increas neutralization of acetic acid. Further addition of NaOH rais steeply due to the presence of free OH ⁻ ions. A typical titrat the model graph.					s are replaced of HCl are new creases slow I raises the c	d by the less utralized. On /ly due the conductance		

•

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	Procedure		Fill a micro burette with the standard N given acid mixture into a clean 100 cm ³ beaker so that the conductivity cell is co Add 0.5 cm ³ NaOH solution from the bu- the conductance. Continue the mea addition of 0.5 cm ³ of NaOH till 10 cm ³ . versus volume of NaOH on X-axis. The c two breaks; the first one corresponds t	aOH solution. Pipette out 50 cm ³ of the beaker. Place the conductivity cell in the ompletely immersed in the acid mixture. Irette. Stir the solution gently and record surement of conductance after each Plot a graph of conductance on Y- axis conductance titration curve is marked by o the equivalence point of HCl (V ₁ cm ³) H (V ₂ cm ³). From the graph, find the NaOH required to neutralize the acids
7	Reaction Equ	ation	NaOH + HCl ——	→ NaCl + H₂O
			NaOH + CH3COOH	← CH ₃ COONa + H ₂ O
	Observation			
	Look-up Output	Table,	Vol. of NaOH (cm³)	Conductance (mS)
			0.0	
			0.5	
		-	1.0	
			1.5	
			2.0	
			2.5	
			3.0	
			3.5	
			4.0	
			4.5	
			5.0	
9	Sample		Normality of NaOH = N (to be	e given)
	Calculations		Volume of NaOH required to neutralize I	$HCl = V_1 cm^3$
			Volume of NaOH required to neutralize (
			$N_{\rm HCl} = \frac{[N \times V]_{\rm NaOH}}{50} = \frac{\dots \times V_1}{50} = \dots$ $\rm NCH_3 COOH = \frac{[N \times (V_2 - V_1)]_{\rm NaOH}}{50} = \dots$	=(b)
				ss of HCl= 'a' x 36.5 = _{DH} x Eq.mass of CH ₃ COOH = 'b' x 6

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10	Graphs, Outputs	
		V_1 V_2
1	Results & Analysis	 Normality of HCl =N Weight of HCl per liter =g Normality of CH₃COOH =N Weight of CH₃COOH per liter =N
12	Application Areas	The experimental determinations of the conducting properties of electrolytic solutions are very important as they can be used to study quantitative behavior of ions in solution.
		They can also be used to determine the many physical quantities such as degree of dissociation and dissociation constants of weak acids and bases, ionic product of water, solubility and solubility
13	Remarks	
	Faculty Signature with Date	

Experiment 03 : Determination of Viscosity co-efficient of the given Organic liquid

-	Experiment No.:	3	Marks		Date Planned		Date Conducted	
1	Title	Deterr viscor		Viscosity	co-efficient	of the giver	n liquid using	g Ostwald's
2	Course Outcomes	Estima metho		efficient of	viscosity of	given organ	ic liquid using	g Ostwald's
3	Aim	Deterr viscor		Viscosity	co-efficient	of the giver	n liquid using	g Ostwald's
4	Material / Equipment Required		10ml grad Organic L	•				
5	Theory	flowin the fo while slow backw betwe the ta	Viscosity arises due to frication between moving layers of a liquid. A liquid lowing through a cylindrical tube of uniform diameter is expected to move in he form of molecular layers. Layer close to the surface is almost stationary while that t the axis of the tube moves faster than any other intermediate layer. A slow moving layer excerts a drag or friction on its nearest moving layer packwards. This property of the liquid, which retards or opposes the motion between the layers, is called viscosity. The Coefficient of viscosity is defined as he tangential force per unit area required maintaining a unit velocity gradient between the two successive layers of the liquid situated unit distance apart. The					

6	Den	SKIT Doc Code:		Teaching	Process		ev No.: 1.0
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Copyright	t ©2017. cAAS	. All rights reserve		cosity of a liquid	is aiven by the	Poiseuille's form	ula.
			$\eta = \frac{\pi \operatorname{pr}^{4} t}{8 \operatorname{Vl}}$ Where 'v' is the pressure betwee volumes of the under identical of $\frac{\eta_{1}}{\eta_{2}} = \frac{t_{1} d_{1}}{t_{2} d_{2}}$ The time't' taken tube is determine the taken taken the taken taken the taken take	volume of the l en the two ends o two different liqu conditions then, n by the given liq ned. The time't' s measured. Kno efficient of viscos	iquid, 'r' is the of the tube is th iids are allowed quid to travel t taken by stanc wing the densi	radius of the tul le Coefficient of v d to flow through hrough a certair lard liquid to tra ties of the two li	be and 'p' is the viscosity. If equal In the same tube In distance in the avel through the quids (d1 and d2)
6 P	Procedur	9	Take a dry viso Immerse the viso cm ³ of the given Suck the liquid mark. Allow the when the level of Remove the viso tiquid from the rinse the viscom Take out the viso average time of Using a thermo temperature.Fro	cometer. (Do not cometer in wate a liquid into the w and fill the bulk e liquid to flow of the liquid cross cometer from the viscometer into the viscometer and f flow for deionize ometer note the myour teacher, g r) and η_2 (Visco coefficient of org	r bath and fix it ider limb of the o on the narrow down through es the lower m e stand. Remov the beaker. Usi over for 20 minu allow a similar d water.(Use a c e temperature get the values c osity coefficient	vertically to a s viscometer usin v limb slightly a the capillary. Sta ark. Note down t ve the rubber tui ng acetone (through thes. procedure for different pipetter of the water b of d1 (density of co of water)	tand. Transfer 15 g a pipette. above the upper art a stop clock the time of flow. be. Pour out the ough a dropper) determining the for water). bath. This is lab
7 M	1odel Dia	agram	$\eta_1 = \frac{t_1 d_1}{t_w d_w} X \eta_v$		$\eta = \frac{\pi}{8}$		milli poise
L	Dbservati .ook-up Dutput		OBSERVATION /	AND CALCULATI		v in seconds	
				Trial 1	Trial 2	Trial 3	Average
			Water				
			Test liquid				
	Sample Calculatio	ons	Lab temperature	9 =	°C		

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		d1 (density of organic liquid) =	g cm ⁻³
		d _w (density of water) = g cm ⁻³	
		η "(Viscosity coefficient of water) =	millipoise
		$\frac{\eta_1}{\eta_w} = \frac{t_1 d_1}{t_w d_w}$	
		$\eta_1 = \frac{t_1 d_1}{t_w d_w} X \eta_w = \frac{\times \times \times}{\dot{c}}$	=milli poise
10	Graphs, Outputs	Viscosity coefficient of the given liquid	
11	Results & Analysis	Viscosity coefficient of the given liquid =	millipoise.
12	Application Areas	Viscosity is how engineers measure the resist	ance of fluids to shear stress.
		The viscosity equation is useful for calculatin know the force being applied to the fluid and t	
13	Remarks		
14	Faculty Signature with Date		

Experiment 04 : Keywords and identifiers

-	Experiment No.:	4	Marks		Date Planned		Date Conducted	
1	Title	Colori	metric estin	hation of Cop	per.	I		
2	Course Outcomes	Comp	ute the amo	ount of Cu by	measuring a	absorbance u	using optical I	method
3	Aim	Colori	metric estin	nation of Cop	per by a give	enCuSO4 sol	ution .	
4	Material /	>	Photo col	orimeter				
	Equipment	\succ	Cuvate tu					
	Required		-	metric flask				
				ılphate solut	ions			
			NH3 solut		<u></u>	<u></u>		
5	Theory, Formula, Principle, Concept	part (
		Thus,	$ _{o} = _{a} + _{r} + _{t}$					
					I			
		Absor	bance is giv	en as A = lo	g $\overline{I_t}$			
			e, E = molar colo C= Mola	red substand		wave length	ny particular 1 of light,	
		Chem wavel wavel the th consta	When the bance, A, ag ical analysis ength is kn ength by a nickness of ant, the abso A series of	e path length lainst concer s through me lown as colo substance in the solution orbance direct f solutions with	atration, c, giv easurements primetry. The solution vari n. When the ctly depends th different c	ves a straight of absorptic absorbance ies directly v thickness upon the cc oncentration	on of light of e of light of vith its conce of the med	a particular a particular ntration and ium is kept nonium ions

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pyri	ight ©2017. cAAS. All rights re	served.		T I		Ť
			on is also meas		ce of cupramm unknown volume	
6	Procedure	five flasks. Take solution to each minutes, set the instrument. The same settings. Draw a on y- axis. (Draw	e the unknown s n one of the six fl e absorbance of en, measure the calibration curve w a straight line p	olution in the si asks. Dilute up to first solution to absorbance of r by volume of C passing through solutions, find o	0, 15 and 20 cm ³ x flasks. Add 5 c o the mark and r zero at 620 nm emaining five sc CuSO₄ on x-axis a the origin). Using out the volume	cm ³ of ammoni nix well. After 1 radiations in th plutions with th and absorbanc g the graph and
	Model Diagram					
7	inodet Blagram					_
8	Observation Tak Look-up Tak Output		Vol. of CuSO₄ in cm³	Volume of ammonia sol. in cm³	Concentration of copper =1.018 mg x vol. of	Absorbance
8	Look-up Tak	ble,	in cm ³	ammonia sol. in cm³	of copper =1.018 mg x	Absorbance
8	Look-up Tak		0.0	ammonia sol. in cm ³	of copper =1.018 mg x vol. of	Absorbance
8	Look-up Tak	ble,	0.0 5.0	ammonia sol. in cm³	of copper =1.018 mg x vol. of	Absorbance
8	Look-up Tak	(Blank sol.)	in cm ³ 0.0 5.0 10.0	ammonia sol. in cm ³ 5.0 5.0 5.0	of copper =1.018 mg x vol. of	Absorbance
8	Look-up Tak	(Blank sol.) (Blank sol.) 1 2 3	in cm ³ 0.0 5.0 10.0 15.0	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of	Absorbance
8	Look-up Tak	(Blank sol.) (Blank sol.) 1 2 3 4	in cm ³ 0.0 5.0 10.0 15.0 20.0	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of	Absorbance
8	Look-up Tak	(Blank sol.) (Blank sol.) 1 2 3	in cm ³ 0.0 5.0 10.0 15.0 20.0 25.0	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of	Absorbance
	Look-up Tat Output	(Blank sol.) (Blank sol.) 1 2 3 4 5 	in cm ³ 0.0 5.0 10.0 15.0 20.0 25.0 Unknown	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of solution	Absorbance
	Look-up Tak	(Blank sol.) (Blank sol.) 1 2 3 4 5 1000 cm ³ of sto 249.54 g of CuS	in cm ³ 0.0 5.0 10.0 15.0 20.0 25.0 Unknown ck solution conta $O_{4}.5H_{2}O = 63.54$ g	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of solution	
	Look-up Tak Output Sample	$(Blank sol.)$ $(Blank sol.)$ 1 2 3 4 5 $1000 cm3 of sto$ $249.54 g of CuSO$ $4 g of CuSO_{4}.51$ solution $1 cm3 of CuSO_{4}.51$	in cm ³ 0.0 5.0 10.0 15.0 20.0 25.0 Unknown ck solution conta $O_4.5H_2O = 63.54$ g $H_2O = 63.54 \times 4$ / $5H_2O = 1.018/100$	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of solution 4. 5H ₂ O g of Cu per 100 of Cu = 1.018 mg	00 cm ³ of stoc
9	Look-up Tat Output Sample Calculations	$(Blank sol.)$ $(Blank sol.)$ 1 2 3 4 5 $1000 cm3 of sto$ $249.54 g of CuSO$ $4 g of CuSO_{4}.51$ solution $1 cm3 of CuSO_{4}.51$	in cm ³ 0.0 5.0 10.0 15.0 20.0 25.0 Unknown ck solution conta $O_4.5H_2O = 63.54$ g $H_2O = 63.54 \times 4$ / $5H_2O = 1.018/100$	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of solution 4. 5H ₂ O g of Cu per 100	00 cm ³ of stoc
9	Look-up Tak Output Sample	$(Blank sol.)$ $(Blank sol.)$ 1 2 3 4 5 $1000 cm3 of sto$ $249.54 g of CuSO$ $4 g of CuSO_{4}.51$ solution $1 cm3 of CuSO_{4}.51$	in cm ³ 0.0 5.0 10.0 15.0 20.0 25.0 Unknown ck solution conta $O_4.5H_2O = 63.54$ g $H_2O = 63.54 \times 4$ / $5H_2O = 1.018/100$	ammonia sol. in cm ³ 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0	of copper =1.018 mg x vol. of solution 4. 5H ₂ O g of Cu per 100 of Cu = 1.018 mg	00 cm ³ of stoc

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11	Results	REPORT : Volume of CuSO ₄ in the unknown solution =cm ³
		Mass of Cu in the unknown solution =mg
12	Application Areas	Colorimeters are widely used to monitor the growth of a bacterial or yea
		culture.
		Colorimeters are used to measure and monitor the color in various foods an
		beverages, including vegetable products and sugar.
4.9		
	Remarks	
14	Faculty Signature	
	with Date	
	with Date	

Experiment 05 : Determination of pKa of the given sample using pH meter.

-	Experiment No.:	5	Marks		Date		Date	
			i la la		Planned		Conducted	
1	Title	Detei	rmination of	f pKa of the g		usina pH me		
2	Course Outcomes			f pKa of the g				
	Aim			f pKa of the g				
-	Material /		Digital Pl					
-	Equipment Required		10ml Bur					
		•	100ml be	eaker				
		•		ng glass elect	rodes			
				id(HCOOH OF)		
		•	1N NaOH	I Solution				
		•	Stirrer					
				olutions(pH4, p				
5	Theory			n acid, which				
				acid CH ₃ COO		make a solu	ition of this a	cid, a part of
				les dissociate	<u>.</u>			
		, °		H ₃ COO ⁻ + H ⁺				i
		Fortr	lis reaction,	the equilibriu	im constant,	ka, is given	by the equal	ion:
		Ka	$[H^{+}]X[C$	H ₃ COO				
		Ки	$=\frac{[H^+]X[C]}{[CH_3]}$	COOH				
			-	n as acid diss	ociation con	stant.		
				garithm to bas			ic., pKa = - loo	g ₁₀ Ka.
		Cons	ider a solut	ion of a weak	k acid; say ad	cetic acid, in	a beaker. Le	et 'Ka' be the
		acid (dissociation	constant.				
				neutralized				
				n of base to t				
				artial neutrali		is related to	o pKa of the	acid by the
				selbalcs equa	ition,			
		ո⊔	- nKa±lo	Salt				
		pu	=pKa+log	Acid				
				e acid agains	st NaOH th	e pH of the	e mixture in	the beaker
	1	1 WC		acia agains				and bound

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		get a 'S' shape equivalence poi the Henderson e PROCEDURE: Pi Immerse the co terminals to a pl micro burette in measure the pH Plot a graph equivalence poi half equivalence	d curve. We fin nt. At half equiv equation pH bec pette out 25 cm ombined glass e H meter. Measur increments of C . (After the jump of ΔpH/ΔV ag nt. Plot another e point (Which is	d that there w valence point, l comes equal to of the given w electrode into t re the pH of the 0.5 cm ³ . After ea in the pH, take ainst volume graph pH/ volu nothing but pk	vill be sharp jum Salt] = [Acid]. Thu pKa at half equiv weak acid into a the acid. Connect acid. Add NaOH ach addition, stir six more reading of NaOH and ume of NaOH, an (a).	us, according to valence point. 100 cm ³ beaker. It the electrode solution from a the solution and gs). determine the d note the pH at		
6 Procedu 7 Model I	Jiagram	pipette. Immerse and connect the burette with the 0.5cm ³ , stir the Continue the pre- few more reading Plot a graph of Plot a graph of	e a glass electro e cell to a pH r base (sodium h solution caref ocedure till the gs after that. Tal apH/aV agair pH (ordinate) a	ode - calomel e neter. Measure nydroxide). Now ully, and meas pH shows a te pulate the read nst V and deter gainst the volu	mine the equivant me of sodium h	bly into the acid acid. Fill a micro le increments of ter 10 seconds. ase rapidly. Take		
8 Observa Look-up Output			P ^H	Δ٧	Δ Ρ ^μ	Δ Ρ ⁺ ΔV		
		0.0						
		0.5						
		1.0						
		1.5						
		2.0						
		2.5						
		3.0						
		3.5						
		4.0						
		4.0			1	1		
		ΛΕ						
		4.5 5.0						

	ght ©2017. CAP Graphs	SKIT Doc Code: Title: AS. All rights reserved		Chemistry Lab	Rev No.: 1.0 Date: 04-08-2019 Page: 18 / 36 Equivalence point (V) alf Equivalence point (V/2) Volume of NaOH in cm3
11	Results		REPORT: The	e pKa of the given acid =	
12	Applicat			ement of pH is used in medical elec	tronics engineering.
13	Remarks	6		· · · · · · · · · · · · · · · · · · ·	
14	Faculty with Dat	Signature e			

Experiment 06 : Flame photometric estimation of sodium and potassium.

-	Experiment No.:	6	Marks		Date Planned	Co	Date Inducted	
1	Title	Flam	e photomet	ric estimation	of sodium a	nd potassium.		
2	Course Outcomes	-		U		ng Flame photo	metric.	
3	Aim	Flam				nd potassium.		
4	Material / Equipment Required		Stock sol 6 numbe	notometer FLA lutions of Na*a ered 100 ml vo pettes: 1, 2, 10 r	and K⁺ , c = 1 lumetric flas	mg/ml.	B • 10	oml urette 20ml
5	Theory	salts	- End of the second sec	CIGITAL DA METER	a vapour, w	which contains n	Flame p is an emission used detection If a containing	technique for the of metals. solution g metallic
		excite by ma these where	electrons fro ed from gro aking use of electrons v e n=2,3,4	f thermal ener will return to t	to higher er gy of flame. he ground s	nergy state (En) From higher en tate by emitting	ergy states	
		NaC			sociation	sion) Na(g) + Cl (g)		
					K*			

	and the second second	SKIT		Teaching Process		Rev No.: 1.0
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Copyri	ight ©2017. cA	AS. All rights reserved	Excitation Er KCl (s)KCl (g) KCl (s)KCl (g) Flame photometer co these elements.It is easily excited (sodium A flame photometer i fuel gases, an automis filter of the element between the flame a oxygen is used as oxi 1900°C. The whole ar rate of introduction o analyte is aspirated ir flame is collected by which permits only investigation into the concentration and nat	orrelates the emitt simple and rapid n and other alkali m is composed of the ser, burner, photos t whose concentrand the detector. F idant. Combination nalysis depends of f the sample and allow the radiation of the radiation of the sample and allow the radiation of photocell. The out ture of the element Vapourisation MX	reed radiations with t method for the ele- netals). e pressure regulato sensitive detector an ration is to be dete Propane gas is use of these two will gi n the flow rate of the droplet size. The sa ugh automiser. Rad owed to pass throuch characteristic of the typut from the photo t Dissociation MX	the concentration of ements that can be r and flow meter for nd output recorder. A ermined is inserted d as fuel and air or ive a temperature of he fuel, oxidant, the mple containing the iation from resulting ugh an optical filter, he element under ocell represents the Thermal excitation M (gas) + Gas
					(g) Flame emission,	h●
6	Procedu		Flame photometer us improperly! Switch the instrument Note: Check the flame the gas valve immedia Transfer 5,10,15,20 an prepared by weighing dissolving the crysta water and mixing. The flasks and dilute up to the suction capillary of read zero. Place each the instrument to rea between each reading and place the solution	t on and off under s e during work if it g ately With Eppend nd 25 cm ³ of stanc g accurately 2.542g Is and diluting the e solution gives 1pp o the mark with dis of the instrument an n of the standard s id 5,10,15,20 and 2g g). Dilute the given	supervision! goes out, close orf flame photomete dard sodium chlorid g NaCl into a 1 liter e solution upto the om /ml) into 100ml stilled water. Place t nd set the instrumer solutions in the sucti 5 respectively (rinse n test solution upto t	er: de solution (which is volumetric flask and e mark with distilled standard volumetric the distilled water in nt to ion capillary and set e with distilled water the mark, shake well

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4		SKIT			aching Proce	SS	Rev No.: 1.0
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	ght ©2017. cAA		calibration c axis). From t given test s contains 23 Determination the same pr 1. Let the ins 2. Feed disti 3. Select the "Elements 4. Turn the c o". Pull the readout to Readjust C 5. Aspirate th (solution n 350 (on up 6. Aspirate c 7. Aspirate s then stan 8. Repeat 3-	he calibration olution and fr g of Na). on of Potassiu ocedure gives trument warr lled water to t e element Na wahl". uter knob "Me e "Kompensat 0. Press the " reading with ne most conc umber 6) and permost scal listilled water candard soluti dards 4, 5, 6. 7 for solutions	curve, find c om which ca in above for s n up for 5-10 the instrument by turning th essbereich" in on I" knob sli Kompensation "Kompensation (Kompensation adjust reado e) using inne – the instrum ions no. 1, 2, 3 Record the reas of potassiun	out the volume of the alculate the amour standard solution of odium. minutes. nt. e selector nto position "10 ghtly out and adjust on I" knob back. ion II" if necessary. ndard solution put to approximatel r "Messbereich" knot nent should read 0. 3, test solution, and esults.	it of Na (58.5 g of Na(f potassium and follo t t
7	Model D	Diagram		Flame	File Lens Oxid Fuel gas To drain		r Amplifier and Readout
	Observa Look-up Output	tion Table, Table,	Volume of sodium chloride solution (cm ³)	Concentrati on of Na = 500 x vol 50 (ppm)	Emission Intensity	Volume of potassium chloride solution (cm ³)	Concentr Emission ation of K Intensity = 500 x vol 50 (ppm)

1		KIT Code: BS-SKIT.Ph		aching Proce	255	Rev No.: 1.0 Date: 04-08-20			
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		2.0	20		2.0	20			
		4.0	40		4.0	40			
		6.0	60		6.0	60			
		8.0	80		8.0	80			
		10.0	100		10.0	100			
		Test			Test solution				
		solution							
	Sample Calculations	DETERMINA	ATION OF SOI	DIUM:					
	Calculations	1ml of NaCl	Weight of Sodium per ml of the solution = 1 mg 1ml of NaCl solution contains 0.002542g of NaCl 58.5 g of NaCl contains 23 g of Na						
		0.002542 g	23 0.002542 g of NaCl contains = × 0.002542 58.5						
		1ml of NaCl Therefore X X ×0.0025 = Therefore th solution	= 1 mg Therefore 1ml of NaCl solution contains 1 mg of Na 1ml of NaCl solution contains 0.002542g of NaCl Therefore Xml of NaCl solution contains = X × 0.002542g of NaCl =×0.002542g of NaCl =						
			Therefore, Y g of NaCl contains						
			23 =×Y =g=mg						
		DETEDMINI	58.5						
		Weight of p 1ml of Kcl so	ETERMINATION OF POTASSIUM: /eight of potassium per ml of the solution = 1 mg ml of Kcl solution contains (0.001909g of KCl 4.5 g of KCl contains 39 g of K						
		1ml of KCl s	39 =×0.001909 =1 mg 74.5 Therefore , 1ml of KCl solution contains 1 mg of K 1ml of KCl solution contains 0.001909g of KCl Therefore, X ml of KCl solution contains = X × 0.001909g of KCl =× 0.001909g of KCl						
		calculated k	by knowing th	of K preser e equivalent	nt in above test so weight of K and mo 39	olution (X ml) can			
		i neretore, Y	g of KCl cont	lai(15 =	-× Y =g 74.5				
	0				=mg				
10	Graphs	Calibratic	n curve						

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Copyri	Title: ght ©2017. cAAS. All rights reserve	Engineering C	nemistry Lab	Page: 22 / 36			
	ght ©2017. cAAS. All rights reserve	e Emission Intensity	Emission Intensity Conc. of Na (ppm)	A Conc. of K (ppm)			
11	Results & Analysis			1.12			
		Result: The weight	ght of Na ⁺ present in the given test sc	olution =mg			
The weight of K+ present in the given test solution=							
12	Application Areas	This method is used in determining in ion concentration in BIOLOGICAL FLUID in medical electronics engineering.					
13	Remarks						
	Faculty Signature with Date						

<u> PART - B</u>

Experiment 01 : Determination of Total hardness of Hard Water sample by using Standard Na2EDTA solution.

-	Experiment No.:	1	Marks		Date Planned		Date Conducted		
1	Title	Deterr	Determination of Total hardness of Hard Water sample.						
2	Course Outcomes		stimation of total hardness of given sample of hard water sample using complexometric titration.						
3	Aim		mination of DTA solutior		ness of Hard	l Water sar	nple by usir	ng Standard	
4	Material / Equipment Required	2. 3. 4. 5. Reage 1. 2. 3. 4.	Pipette Conical fla F annel ents Na2EDTA Ammonia Hard wate	ask Solution solutions er Solution Cl Buffer sol	ution				
5	Principle	Hardn salts in of ca sulpha (EDTA	ess of wate n it. Total ha lcium and ates etc., of) is a reager	er is mainly o Irdness is the Magnesium Calcium and It, which read	sum of tem) and perm d Magnesium cts with meta	porary hardr anent hardı n). Ethylene ıl ions like Ca	calcium and ness (due to k ness (due to diamine tetra a ²⁺ &Mg ²⁺ form ermine the co	oicarbonates o chlorides, a acetic acid ing complex	

6	Sector Sector	SKIT			Tead	ching Pr	ocess		ev No.: 1.0		
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			of hardn		sing substa	ances. CE-JCCOI		C	C-COD4		
			-	4—1 <u>4</u>	╤┓╔╋		ſ	4−ŀł͡C−Œ			
				C		CF-CCO		C	CE-ICODH		
6	Procedu	7	Eriochro relatively are wine- wine-rec metal io be repre M ²⁺ Indicator Observe is sensit around 1 buffer is	The completion of the reaction (end point of the titration) is identified using Eriochrome black- T indicator. This is an organic dye, blue in colour. It also forms relatively less stable complexes with bivalent metal ion of Ca &Mg etc., which are wine red in colour. Therefore addition of the indicator to hard water produces wine-red Colour. When EDTA is added to hard water, it first reacts with free metal ions and then attacks the metal-indicator complex .The latter reaction can be represented as M^{2*} Indicator complex + EDTA $\rightarrow M^{2*}$ EDTA complex (COLOURLESS) +free Indicator (Blue) so at the end point a change from wine red to blue colour is Observed. Since the reaction involves the liberation of H* ions and the indicator is sensitive to the concentration of H* lons (pH) of the solution a constant Ph of around 10 has to be maintained. For this purpose ammonia-ammonium chloride buffer is used. Part-A: Preparation of standard EDTA solution							
0	 Procedure Part-A: Preparation of standard EDTA solution Weigh the weighing bottle containing disodium salt of Na, accurately and transfer the salt in to the funnel placed on a 250 cm³ volur flask. Weigh the bottle again .The difference between the two weights will the amount of Na₂EDTA transferred. Pour Small quantities of water over the on the funnel and transfer the salt in to the Flask. Wash the funnel with the water 3-4 times; Dissolve the salt by adding 5ml 1:1 Ammonia and make u solution to the mark and shake well for uniform Concentration Part-B: Estimation of hardness of water Pipette out 25 cm³ of the given sample of hard water in to a clean of flask .Add 5 ml of NH₃-NH₄Cl buffer followed by 3-4 drops of Eriochrome bindicator .Titrate this against Na₂EDTA taken in a burette till the colour charform wine red to pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the solution of the pure blue .Note down the burette reading and repeating the pure blue .Note down the burette reading and repeating the pure solution approximate and the pure solution to the pure solution the pure solution the pure solution the pure solutio						to a clean conical iochrome black T e colour changes				
	Block,	Circuit	- J	<u> </u>	oncordant						
	Expected						NIL				
	Observal Look-up Output	tion Table Table			Trial I	Tr	ial 2	Trial 3	Indicator and colour change		
			Final reading	burette)							
		Initial Readin	burette g					EBT indicator Wine red to			
			Volume EDTA run de cm ³	e of own in					clear blue		
	Sample		OBSERV	ATION A		ULATION	N:				
	Calculati	ons			tion of Na₂						
			Weight	of the w	eighing bo	ottle +Na	2EDTA = W1=		g		

	SKIT		Taaabir							
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		Weight of t	he weigl	hing bottle	⊖ = W₂=	g				
		Weight of the Na₂EDT	2)= g							
		Molarity of EDTA	Molarity of EDTA solution = $\frac{\text{Weight of Na}_2\text{EDT}(W_1-W_2)X4}{\text{Gram molecular wt. of Na}_2\text{EDTA}} = \frac{372}{372}$							
		PART-B : Estimatio			innoiceu	= M (<i>a</i>)				
		EDTA in burette	Trial I	Trial 2	Trial 3	Indicator and colour change				
		Final burette reading								
		Initial burette Reading				EBT indicator — Wine red to clear blue				
		Volume of EDTA								
		run down in cm ³								
		Volume of Na₂EDTA us 1000cm³of 1M EDTA		n ³ .00 g of Ca	aCO ₃					
		Therefore <i>b</i> cm³of <i>a</i> m	$b \text{ cm}^3 \text{ of } a \text{ molar EDTA} = \frac{bXaX100}{1000} = \dots (c) \text{ g of CaCO}_3$							
			25 cm ³ of hard water contains =(c) g of CaCO ₃ Therefore 10^{6} cm ³ of hard water contains							
			=	$\frac{c \times 10^{\circ}}{25}$	=	——— ppm				
		Total hardness of Wat		=pp	m of CaC	O ₃				
	Outputs	Total hardness of Wate		=ppr						
	Results & Analysi									
12	Application Areas	 Complexomtri hardness of wa 		on is an e	fficient m	nethod for determining level of				
13	Remarks		-							
	Faculty Signatu with Date	Ire								

Experiment 02 : DETERMINATION OF CALCIUM OXIDE IN CEMENT SOLUTION.

-	Experiment No.:	2	Marks		Date Planned	Date Conducted		
-		DET						
1	Title					MENT SOLUTION		
2	Course Outcomes	Calc	ulate % of Ca	ao in a given d	cement sample	e using rapid EDTA metł	nod.	
3	Aim	DET	ERMINATION	N OF CALCI	UM OXIDE IN	I CEMENT SOLUTION	BY	USING
		STAI	NDARD Na2	EDTA SOLUTI	ON.			
4	Material /	App:	aratus					
	Equipment	6	6. Volumetr	ric flask				
	Required	7	7. Burette					
		8	3. Pipette					
		9	9. Conical fl	ask				
		1	.o. Fannel					
		Reag	<u>gents</u>					
			-					
			1. Concentr	ated Hcl				

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Copyri	ght ©2017. cAAS. All rights reserve	 a. Na2EDTA Solution 3. Cement solution 4. Glycerol Solution 5. Diethyl amine Solution 6. 4N NaOH Solution 7. Patton and Reeder's indicator 	
5	Principle	The major constituents of Portland cement are Silicat aluminum and iron with a small quantity of oxides of composition of Portland cement is as follows CaCO ₃ -63.80%; SiO ₂ -20.7%; Al ₂ O ₃ -5.6% MgO - 3.75%; TiO ₂ -0.23% ; Na ₂ O - 0.21%; SO ³⁺ - 1.75% Use of Eriochrome black-T as indicator gives the tota Mg ²⁺ ions, While Patton & Reeder's indicator would Calcium ions in the presence of Magnesium ions. Fo has to be maintained. Additions of Diethylamine & So purpose.	² alkali metals The average % ; Fe₂O₃ - 2.5%; K₂O - 0.51 %; al concentration of Ca²⁺and d allow estimation of only or this purpose P ^H of 12-14
6	Procedure	 Part A: Preparation of solution of Disodium salt of N Weigh the given disodium salt of Na₂EDTA and trans on a 250 cm³ volumetric flask. Dissolve by adding s Make it up to the mark and shake well to get uniform Part B: Estimation of CaO Pipette out 25 cm³ of given cement solution into a clest com³ of diethyl amine and 5 cm³ of 1:1 glycerol. Adjust adding 10 cm³ of 4N sodium hydroxide solution. Reeder's indicator. Titrate the solution against ED burette until the colour changes from wine red to bly reading and repeat the titration to get concordant value. 	fer on to the funnel placed small amount of DM water. concentration. ean conical flask using. Add st the pH of the solution by Add a pinch of Patton & DTA solution taken in the ue. Note down the burette
	Block, Circuit Model Diagram Reaction Equation Expected Graph	NIL	
8	Observation Table Look-up Table Output		3 Indicator and colour change Patton and Reeder's indicator Wine red to clear blue
9	Sample Calculations		3
		Weight of the Na₂EDTAsalt transferred= g]

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Соруп	ght ©2017. cAAS. All rights reserve	Molarity of Na₂E Weight of Na	a2EDTA salt 2 11ar weight of			M (a)			
		EDTA in burette	Trial I	Trial 2	Trial 3	Indicator and colour change			
		Final burette reading							
		Initial burette reading				Patton and Reeder's indicator			
		Volume of EDTA run down in cm ³				Wine red to clear blue			
		Weight of cement sample in 25 cm ³ = 0.09 = W g Volume of EDTA required to react with 25.0 cm ³ of the cement solution = 1000 cm ³ of 1M EDTA = 56.08 g CaO (Molecular mass of CaO = 56.08) b cm ³ of a M EDTA = $\frac{56.08 \times a \times b}{1000 \times 1}$ g of CaO =							
		$25.0 \text{ cm}^3 \text{ of ce}$			'c 'g of CaO				
		Percentage of (CaO in the ceme	t sample = $\frac{c}{W}$	× 100_=				
				=					
	Outputs	Percentage of C							
	Results & Analysis	REPORT: Percer	ntage of CaO in t	he cement sa	mple =	· · · · · ·			
	Application Areas	This technique engineering.	is applicable	to determine	the quality o	of cement in civil			
	Remarks								
14	Faculty Signature with Date	•							

Experiment 03 : DETERMINATION OF PERCENTAGE OF COPPER IN BRASS

-	Experiment No.:	3	Marks	Date	Date	

State of the second sec	SKIT	Teaching Process	Rev No.: 1.0					
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1	Title	Planned Conducted DETERMINATION OF PERCENTAGE OF COPPER IN BRASS					
	Course Outcomes	Estimation of percentage of Copper in a given alloy by iodometric method.					
3	Aim	DETERMINATION OF PERCENTAGE OF COPPER IN BRASS BY USING STANDARD Na2S2O3 solution.					
	Material / Equipment Required	Apparatus 1. Volumetric flask 2. Burette 3. Pipette 4. conical flask 5. F annel					
		 Reagents Concentrated glacial acetic acid Standard sodium thiosulphate solution (0.025N) Potassium iodide NH4OH Solution Starch indicator Brass solution 					
5	Principle	The chief constituents of brass alloy are copper and zinc. It also contains small quantities s tin, lead and iron. The percentage composition of typical brass is copper 50-90, zinc: 20-40, Tin; 0.6, Lead; 0.2, Iron; 0.1 A solution of brass is made by dissolution of the sample in nitric acid. Boiling with urea destroys oxides of nitrogen. Adding ammonia neutralizes excess acid. The solution is changed to weak acidic medium by adding acetic acid. Potassium iodide is added. Iodine is liberated by the cupric ions. Then the solution is tittered against sodium thiosulphate solution using starch as indicator. The amount of sodium thiosulphate consumed is the measure of the amount of copper present					
6	Procedure	PART A: Preparation of Brass solution: Weigh exactly the given sample of brass into a clean 250 cm ³ conical flask. Add 3cm ³ of 1:1 nitric acid and boil. Add 2 test tube of Dm water and about 1 g of urea. Boil for about 2 minutes destroy oxides nitrogen. Cool the mixture.					
		PART –B: estimation of copper in brass solution. Add 1 test tube of Demineralised water to the solution obtained in part A. Add Ammonium hydroxide drop by drop until a pale blue precipitate is obtained. Dissolve the precipitate by adding 5cm3 of acetic acid and 10cm ³ of 20% KI solution.Titrate the librated iodine against standard sodium thiosulphate solution taken in the burette until the solution becomes PALE YELLOW. Add about 2 cm ³ of freshly prepared starch solution as indicator. Continue the titration by adding sodium thiosulphate solutionStrictly drop by drop until the dark blue coloration disappears, leaving behind white ppt. Repeat PART A and Part B to conduct a duplicate. Calculate the percentage of copper present in brass sample.					
	Reaction Equation	$2Cu^{2^{+}} + 4KI Cu_2I_2 + 4K^{+} + I_2$ $2Na_2S_2O_3 + I_2I + Na_2S_4O_6$					
	Observation Table Look-up Table Output	Burette Sample-I Sample-II Sample-III Indicator and					

Sod. colour							
Copyright ©2017. cAAS. All rights reserved. Final Starch Initial Disappeara Volume of Sod. colour							
Final Starch Initial Disappeara Volume of Sod. Colour							
Initialsolution.VolumeofSod.colour							
Volume of Sod. colour							
Sod. colour	lue						
Thiosulphate							
run down (in cm ³)							
9 Sample OBSERVATION AND CALCULATION: Calculations SAMPLES Sample-1 Sample-2 Sample-3							
Weight of the brass g transferred g	g						
PART –B: Estimation of copper in brass solution.							
Burette readings Sample-I Sample-II Indicator and concernance	lour						
Final Starch solution.							
Initial Disappearance	of						
Volume of Sod.							
Thiosulphate run down (in cm ³)							
SAMPLE 1: Normality of Sodium. Thiosulphate =(a) N Volume of the Sod. Thiosulphate =(b) cm ³ 1000 cm ³ of sod. thiosulphate = 63.54 g of copper $\frac{63.54X \text{ bX } a}{1000} = \frac{63.54 \text{ X} \text{ X}}{1000}$							
$= \dots \dots$							
Therefore, 100g of brass contains = W g of copper	Therefore, 100g of brass contains = W						
(Note : Similarly do the calculation for II and III trial)							
10 Outputs Percentage of copper in brass sample =							
11 Results & Analysis Percentage of copper in brass sample =							
12 Application Areas This method is used to determine composition of metals in an alloys.13 Remarks							
13 Remarks 14 Faculty Signature with Date							

Experiment 04 : DETERMINATION OF PERCENTAGE OF IRON IN HAEMATITE ORE SOLUTION

-	Experiment No.:	4	Marks	Date Planned		Date Conducted	
1	Title	DETI	ERMINATION	I OF PERCENTAGE OF IR	ON IN HAEM	ATITE ORE	

1	SKIT		Teaching Process	Rev No.: 1.0
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	T		Calculate % of Fe in a given ore solution using exter	nal indicator method.
3	Aim		DETERMINATION OF PERCENTAGE OF IRON IN H. BY USING STANDARD K2Cr2O7 SOLUTION.	AEMATITE ORE SOLUTION
	Materia Equipm Require	ent d	Apparatus 11. Volumetric flask 12. Burette 13. Pipette 14. Conical flask 15. Funnel Reagents 1. Concentrated HCl 2. Haematite ore solution 3. SnCl2 Solution 4. HgCl2 Solution 5. Potassium dichromate 6. [K ₃ (Fe(CN) ₆](external)	
5	Principl		Haematite is an important ore of iron containing mair Estimation of involves the dissolution of the ore in I the Ferric (Fe ³⁺) ions in the solution to Ferrous (Fe ²⁺⁾ ike Stannous chloride and the estimation of ferrous against an Oxidizing agent like Potassium dichromation	Hydrochloric acid, reducing ions using a reducing agent ions so obtained by titrating
6	Procedu		 Part A - Preparation of Potassium Dichromate solut Weigh accurately the given potassium dichromate the funnel placed on a 250 cm³ volumetric flask. quantities of DM water and make upto mark. S concentration. Part B Estimation of Iron: Pipette out 25 cm³ of the given Haematite solution Add 5 cm³ of concentrated Hydrochloric acid. H poiling. Add Stannous chloride drop by drop to f solution becomes Colureless. Add 2-3 drops of st Cool the solution to room temperature. Add 2 test tu 	crystals and transfer on to Dissolve by adding small hake well to get uniform in to a clean conical flask. eat the solution nearly to the HOT solution until the annous chloride in excess. be of DM water followed by
			5 cm ³ of Mercuric Chloride at a strech. A silky White the contents of the flask and repeat The experie GREYISH ppt is formed. Titrate the solution ac dichromate solution taken in the burette using po EXTERNAL INDICATOR. In the beginning take out a c using a clean glass rod and mix it with a drop of t paraffin paper. The colour of the drop of indicator of drop of the reaction mixture after every addition of fresh drop of the indicator, appearance of blue or the END point is not reached. At the end point a d fails to give either blue or green coloration. Note do repeat the experiment for agreeing values.	ment if NO PRECIPATE or gainst standard potassium stassium ferricyanide as an drop of the reaction mixture he indicator arranged on a changes to blue. Take out a of K ₂ Cr ₂ O ₇ and mix it with a green colour indicates that rop of the reaction mixture
7	Reactio	n Equation	2FeCl ₃ + SnCl ₂ → 2FeCl ₂ + SnCl ₂ Yellow Colorless	
			SnCl₂ + 2HgCl₂ → SnCl₄ + Hg₂(Silky	Cl ₂ white
			$K_2Cr_2O_7 + 8 HCl \rightarrow 2KCl + 2 CrC$ $(2FeCl_2 + 2 HCl + [O] \rightarrow 2 FeC$ $K_2Cr_2O_7 + 14 HCl + 6FeCl_2 \rightarrow 2KCl$	$l_3 + H_2OX_3$

	AND DE LE COLORIZACIÓN DE LE COL	SKIT	Te	eaching Pro	ocess		Re	v No.: 1	0
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8	Observatic Look-up Output			Trail I	Trail II	Trail III	Indicator change	and	colour
	output		Final burette reading				[K ₃ (Fe(CN)		ernal)
			Initial burette reading				Ŭ		
			Volume of K ₂ Cr ₂ O ₇ run down (in cm ³)				Blue to nc colour of i	-	
9	Sample		PART A: Preparation of	ootassium	dichrom	ate solut	ion		
	Calculatio	าร	Weight of the weighing	bottle + K ₂	$cr_{2}O_{7} =$		g		
			Weight of the weighing	bottle	=		g		
			Weight of the K ₂ Cr ₂ O ₇ sa	alt transfe	rred =		g		
			Normality of K ₂ Cr ₂ C	P_7 solution	$h = \frac{Wt}{Cram}$	of $K_2 Cr_2$	$_{2}O_{7} X 4$. of $K_{2}Cr_{2}$	=	$\frac{X}{49,06}$
			Part B: Estimation of Iro	n [.]	Gran	i Eq. wi	$\cdot 01 \text{ K}_2 \text{ Cr}_2$	$_{2}O_{7}$	49.06
			Burette readings	Trail I	Trail II	Trail III	Indicator change	and	colour
			Final burette reading						
			Initial burette reading				[K ₃ (Fe(CN)		
			Volume of K₂Cr₂O7 run down (in cm³)				Blue to no colour of i		-
			Volume of K2Cr2O7 consu Weight of haematite ore 1000 cm3 of 1NK2Cr2O7	dissolved = 1 equ		iron	solution = 1.	025 g	
			Therefore (b) 55.85 X b X a = 5	cm ³	of		normal	K₂Cr₂	O ₇
				1000					
			25 cm³ of haematite ore 250 cm³ of haematite ore		contains (n	(d) g) of iron
			Therefore, 100g of hae	matite ore	e contain	$\frac{\mathrm{d}x}{1}$	$\frac{100}{025} = -$	X	100 25
			Percentage of iron in giv	on haoma	tito oro or	amplo -			
10	Outputs		Percentage of iron in giv						
	Results & /	Analysis	REPORT : Percentage of						
12	Applicatio	n Areas	This method is used metallurgical process.	to deterr	nine cor	nposition	of metal	s in it	s ore in
	Remarks								
	Faculty with Date	Signature							

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Experiment 05 : DETERMINATION OF CHEMICAL OXYGEN DEMAND (COD) OF WATER

-	Experiment No.:	5	Marks		Date		Date		
					Planned		Conducted		
	Title					DEMAND (CC			
2	Course Outcomes		stimation of total oxidizable impurities present in sewage water through redox ration.						
3	Aim		TERMINATION OF CHEMICAL OXYGEN DEMAND (COD) OF INDUSTRIAL AST WATER SAMPLE BY USING STANDARD FAS SOLUTION.						
4	Material / Equipment Required	1 1 1 2 Reac		ask ated H2SO4 mmonium su n dichromate dicator					
5	Principle	that of impo comp other comp is ad	can be oxidiz ortant parame oounds, aror r oxdisable n oounds, acet	eed by a stror eter in indust natic hydroca naterial are p ic acid etc. a atalyst. Addi	ng oxidizing a rial wastewa arbons, strai present as in re oxidisabe	agent. Chemi ater treatmen ght chain alc apurities in w more effectiv	cal oxygen c t. Straight ch ohol, acids, astewater. S /ely when si	ole materials demand is an hain aliphatic pyridine and straight chain lver sulphate help avoid	
6	Procedure	Weig using wate conc	nh accurately g a funnel. Ac	/ the given F dd 30cm³ of (, make it (FAS and trar dilute sulphu	iric acid follo [,]	250 cm ³ st wed by abou	solution: tandard flask ut 100 cm ³ of for uniform	

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			Add 10 cm ³ of 1:: and 3 drops fer colour changes repeat the titration Part-C: Back titr Pipette out 25 c cm ³ of standard 1:1 sulphuric acid the flask constat temperature. Act taken in the bure	a sulphuric acid roin indicator. T from blue green on to get conco ration: cm ³ of given sa potassium dicl d containing me antly. Reflux the dd 3-4 drops fe ette until the co	containing me Titrate against in to reddish br ordant values. mple of waste hromate soluti ercuric sulphate content of fl erroin indicato lour changes f	ercuric sulphate FAS taken in th own. Note the b ewater into a co on using a pipe e and silver sulp ask for 30 minu r and Titrate ag rom bluish green	Task-using pipette. and silver sulphate e burette until the purette reading and nical flask. Add 25 tte. Add 10 cm3 of hate while shaking utes. Cool to room jainst FAS solution n to reddish brown. concordant values.		
		n Equation							
8	Observa Look-up Output	tion Table Table		Trail I	Trail II	Trail III	Indicator and colour change		
			Final burette reading				Ferroin — indicator		
			Initial burette reading				Blue green to — Reddish		
			Volume of FAS run down (in cm ³)				brown		
9	Sample		OBSERVATION A						
	Calculat	ions		RT A: Preparation of Ferrous ammonium sulphate (FAS) solution: eight of the weighing bottle + FAS = g					
						g			
			Weight of the w		=	g			
			Weight of the F	AS salt transferr	'ed =	g			
			Volume of FAS of	of FAS $=$ 392 consumed in th		solu N(a) n = (b) cm ³			
			Part-B: Back titr Burette	ration: Trail I	Trail II	Trail III	Indicator and		
			readings	naiti			colour change		
			Final burette reading				Ferroin indicator		
			Initial burette reading				Blue green to Reddish		
			Volume of FAS run down (in cm ³)				brown		
			Back titrate valve Amount of pota sample = _ (b)	ssium dichroma	:) cm³ ate (in terms o	f FAS) that has	reacted with water		
			1000 cm ³ of 1N F	AS solution = 1	equivalent of c	xygen = 8 g of o	xygen.		

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		b - c cm ³ of ' a' N FAS solution = $\frac{(b-c) X a X 8}{1000} = \frac{1000}{1000} = \frac{1000}{1000}$
		25 cm³ of wastewater requires (d) g of oxygen
		<u>d x1000</u>
		Therefore, 1000 cm ³ of waste water requires = 25 =
		COD of the given sample of water =mg/dm³ of oxygen_
10	Outputs	COD of the given sample of water =mg/dm³ of oxygen
11	Results & Analysis	REPORT : COD of the given sample of water =mg/dm ³ of oxygen
12	Application Areas	This technique is used to maintain standard parameters in industrial waste
		water in environmental engineering.
13	Remarks	
14	Faculty Signature with Date	

Experiment 06 : Estimation of percentage of available chlorine in the given sample of bleaching powder

-	Experiment No.:	6	Marks		Date Planned		Date Conducted		
1	Title	Estim powd	timation of percentage of available chlorine in the given sample of bleaching						
2	Course Outcomes		timation of % of chlorine in a given bleaching powder sample by lodometric ethod.						
3	Aim				vailable chlori 2S2O3 Solutio		ven sample	of bleaching	
4	Material / Equipment Required	V V. Reage V V V	Mortar ar Volumetr Burette Filenmey ents (I. Concentr (II. Standard (III. Potassiu K. Starch ind	ric flask ver flask. rated glacial a sodium thios m iodide	sulphate soluti	on (0.025N)			
5	Principle				nly used as a e educed with ti				

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		r C C	must Chlorine will libe or less. The ioc	er required, the amo be rate free iodine from line liberated, which ed with standard soc	potassiu 1 is equ	found m iodide s ivalent to	solution the ar	out. when its pH is 8 mount of active
6	Procedure		flask, (This can mortar ar 2. Place 5 potassiur prepared 3. Titrate w colour is 4. Add 1mL 5. Note dow 6. Take a vo 7. Add 5 ml 8. If blue co until the f 9. Record th 10. If no blue colour ap 11. Then, titr colour di added (A solution	1g bleaching powde and be done by first mal nd pestle.) mL acetic acid in m iodide crystals. Po l above and mix with ith 0.025 N sodium obtained. (Deep yello of starch solution and on the volume of sodium of distilled wate acetic acid, 1g potas plour occurs, titrate w olue colour disappea ne volume of sodium e colour occurs, titrate pears. Note down the rate with 0.025 N so sappears. Record the A ₃). Note down the of and sodium	stopper king a par an Erler bur 25 n a stirring thiosulp bw chang d titrate of ium thiosulp with 0.02 rs. thiosulp a volume odium the odium the odium the odium the odium the solume difference	aste of the nmeyer fla nL of blea nod. hate solut ges to pale until the bl sulphate s ponding to dide and 1 5 N sodiu hate solut .025 N iod e of iodine hiosulphat e of sodiu e betweet	the bleach ask and aching tion uni- e yellow lue colo olution o the sa mL star m thios ion add ine solut (A ₂). e solut m thios n the v as	container. ing powder with d add about 1g powder solution til a pale yellow bur disappears. added (V_1). imple used. rch solution. ulphate solution ed (A_1). ution until a blue ion till the blue ulphate solution olume of iodine $A_4(A_4=A_2-A_3)$.
7	Reaction Equat	ion	$A_{4}(A_{4}=A_{2}-A_{3}).$					
8	Observation T	Table, Table,	Bleaching pow	der solution x Standa				
			Trail no.	Volume of ble Powder solution(mL		Burette re Initial	Final	Volume of tit rant(mL)
			Distilled water × Trail no.	Standard sodium thi Volume of ble Powder solution(mL	eaching	te solution Burette re Initial		N) Volume of tit rant(mL)

- /	SKIT		Teaching Process	Re	Rev No.: 1.0			
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		Distilled water >	<pre> Standard iodine solution (0.0 </pre>) 25N)	1	<u> </u>		
		Trail no.	Volume of bleaching		ading	Volume of tit		
			Powder solution(mL)	Initial	Final	rant(mL)		
				million	1 mat			
9	Sample							
	Calculations $(V - A_1) \text{ or } (V + A_4) \times N \times 35.46$							
		mg of Cl ₂ /mL (B) =						
		mL of bleaching powder solution taken						
		1000 mL of bleaching powder solution contains 1000 x B mg of Cl_2						
		ie 1000 mable	aching powder contains 1000	B ma of C	1			
		1.e., 1000 mg bie		b mg or c	2			
		therefore, 100 m	g of 1000 X	В				
		bleaching powd	er contains =					
				10				
				10				
		% of chlorine	available =					
10	Outputs	Available chlorin	e in the given bleaching pow	der is%				
	Results & Analysis		ne in the given bleaching pov					
	Application Areas	This technique i	s used to determine the quali	ity of blead	ching po	owder sample.		
-	Remarks							
	Faculty Signature with Date							

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