A COMPARATIVE STUDY OF COPPER, ZINC AND ALUMINIUM AS ELECTRODES IN THE PREPARATION OF STARCH-BASED POLYELECTROLYTE

ABSTRACT

7 Alum $(Al_2 (SO_4)_3)$. 18H₂O) has been the coagulant mostly used in water treatment process. Alum imported into the country gets stuck at the port because of the congestion of the 8 port. When alum is used to coagulate water, it depresses the pH of the water, thus, 9 requiring a pH booster. Because of the high dependence on alum, alternative coagulants 10 produced with local raw materials will be of great advantage. A starch-based 11 polyelectrolyte was prepared by dispersing 3 g of cassava starch and 10 g of heaping 12 teaspoon of high-test granular calcium hypochlorite (commercially available as HTH) in 13 14 100 ml distilled water; making it up to a litre solution with hot distilled water (temperature 15 80°C – 100°C) and electrolysing by means of stabilized power supply using aluminium rods as electrodes to produce polyelectrolyte A. Copper rods and zinc rods were 16 similarly used to polyelectrolyte B and polyelectrolyte C respectively. To test the effect(s) 17 and interaction of the charging variables, i.e. separating distance, power supply and time 18 of charging, a three-factor experiment (2³ factorial design) was carried out. The variables 19 20 were operated at High Level (+) and Low Level (-). The analysis indicate that the power of charging has the most significant effect on the coagulation characteristics of the starch-21 based polyelectrolyte. Polyelectrolytes A and C were tested with raw water drawn from 22 Opa dam and were found to be effective water coagulants. Polyelectrolyte B on the other 23 hand, adds to the turbidity of the raw water with which it was tested. Polyelectrolyte A 24 was found to be as effective as Alum at natural water condition (pH 7) causing better 25 than 88% turbidity removal with dosage of 65 mg/l. 26

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Keywords: Comparative, Starch-based polyelectrolyte, alum, coagulant, surface water
 treatment.

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31 **1. INTRODUCTION**

32 Water which is absolutely pure is not found in nature, every water vapour condensing in air contains solid

and dissolved gases. As condensed water falls it sweeps up other material from the air and becomes

34 contaminated on reaching the ground, running over the surface and percolating through the various strata

of the soil [1]. These contaminants may be removed by passage through the soil as a result of filtration

36 and exchange and adsorption reactions; some may be removed from the surface water by sedimentation

- 37 and biological activity; others may be removed by specific engineered process in water treatment plants.
- 38 New impurities are introduced by dissolution and exchange reactions in the soil, others by accumulation
- 39 of decomposition and reactions with the materials of the distribution system [2]. In order to provide a
- 40 potable water supply, raw water from the source has to be treated. The source may be a river, reservoir,
- 41 well or natural lake. Each of these sources have their own peculiar impurities and means of removing

them. A large portion of the suspended particles in water are sufficiently small that their removal in the sedimentation tank is impossible at reasonable surface overflow rate [3]. Coagulation is also called chemically assisted sedimentation; it is the process of causing the finely divided particles in water to cluster together and form larger particles (flocs) and later settle before passing the water to the filter [4]. The four major mechanisms of coagulation are double-layer compression, adsorption and charge neutralisation, enmeshment in a precipitate and adsorption by polymers, and inter-particle bridging. The understanding of the four processes is necessary for understanding the phenomena of coagulation [5].

49 The coagulation unit is essential in removal of impurities and alum (aluminium sulphate) is the most 50 common coagulating agent. There is a need for a local substitute due to the rapid increase in the number 51 of water treatment plant and the constantly increasing cost of alum. Cationic starches have long been 52 used to flocculate negatively charged particles from aqueous suspension. Safi et al. [6] reported the 53 preparation of 2-aminoethyl ether derivatives of starch which readily flocculated aqueous suspensions of 54 negative colloids. Salim et al.[7] reported the use of high molecular weight polyamines, polyalkylamines 55 and polyacrylonitriles as flocculating materials. Cationic quaternary ammonium starch ether which was 56 found to be an excellent flocculant for materials in aqueous system [8].

57 Though these starches are effective flocculating agents for aqueous suspension of negatively charged 58 inorganic and organic particle. Their precise modes of operation were not precisely understood. Sirin [9] 59 was able to give a simple explanation, the simple ionic attraction between the product formed and the 60 colloidal particles leading to the production flocculants - particle aggregates large enough to settle from 61 suspension; the polyelectrolyte, according to his discovery also form bridge structure that leads to 62 precipitation. Tenney et al. [10] explained that many of the suspended particles in raw water are too small to settle out by themselves in a separation by sedimentation process. The benefit of adding of polymer is 63 64 derived from the agglomeration of these individual particles into larger clumps or flocs allowing them to 65 settle and be removed. The list of synthetic polymers which have been successfully applied to treatment 66 of water.

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68 2. MATERIALS AND METHODS

Locally purchased cassava was peeled and washed. They were then crushed, with a cassava grater machine, then oaked for some time in water after which they were sieved. The filtrate was allowed to settle for some hours and the settled starch was separated from the water by careful decantation of the top water. The residue was then air dried, grounded and stored in a desiccator to prevent the adsorption of water vapour from the atmosphere. The material obtained from this process was then used in the preparation of the starch-based polyelectrolyte.

- 75 2.1. Preparation of Starch-based Polyelectrolyte
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77 Tablets of calcium hypochlorite (Ca(OCI)₂), commercially available as HTH, were ground and stored in a 78 desiccator [11]. Three gram of starch and 10 g of hypochlorite were made into a well-mixed slurry with 79 100 ml distilled water in a one-litre beaker. Hot distilled water (70°C - 100°C) was added to the slurry, 80 making it up to a litre solution with continuous stirring. The resulting solution was electrolysed on a 81 stabilized power supply with aluminium rods, copper rods and zinc rods as electrodes one after the other, to produce polyelectrolytes A, B and C respectively. The solution was gently stirred in each case 82 throughout the process of charging. The variables i.e. power, P, separating distance, D and time of 83 charging, T; were tested at the high level (+) and low level (-) for each of the electrodes to generate a 2³ 84 85 factorial runs for each of the electrodes (Tables 1 and 2).

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87 Table 1: Electrolysing Variables Values at High and Low Levels

	Variable	$\mathbf{\vee}$	Hig	gh Level (·	+)	I	Low Lev	el (-)	
	Power product of voltage and	d current (watts)		400			100		
	Separating distance	e (mm)		150			50		
	Time of charging	(min)		60			30		
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89									
90									
91	Table 2: Factorial De	sign Table							
	Run	1	2	3	4	5	6	7	8
	Power, P (Watts)	-	+	-	+	-	+	-	+

Separating Distance, D (mm)	-	-	+	+	-	-	+	+
Time, T (mins)	-	-	-	-	+	+	+	+

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93 2.2. Coagulation of Water Samples

Raw water from the university dam, Opa Dam, was collected and used in testing the effectiveness of the
starch-based polyelectrolyte as a coagulant under neutral condition i.e. natural water, pH 7.

96 Treatment in each case consisted of dosing varying amount of the prepared polyelectrolyte into the water,

97 mixing rapidly for 60 seconds and then slowly for 10 minutes in a jar test stirrer. The water was then
98 allowed to settle for 30 minutes. Coagulation effectiveness was determined by measuring turbidity of the

99 water sample before and after treatment using a Hach's Model 2100A turbidimeter.

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102 3. RESULTS AND DISCUSSION

Preliminary coagulation test carried out with the starch-based polyelectrolyte as coagulant revealed the following: the ratio of hypochlorite to starch must not be less than 3; the polyelectrolyte can be stored at room temperature for more than 12 days without materially losing its effectiveness as a coagulant; a better turbidity removal can be achieved if the settling time is increased; and when water at natural pH and temperature was used to make the slurry to a litre solution without stirring, the turbidity removal is relatively low compared with when hot distilled water is used with continuous agitation [12].

3.1. Comparing the Effectiveness of the Polyelectrolytes

Varying amounts, in millilitres, of the polyelectrolytes were used to treat litres of raw water sample (turbidity, 28 NTU). Table 3 shows the conversion from millilitre to milligrams per litre; the conversion is on the basis that 1 mg/l of the solution prepared as indicated in the preceding contain 3 mg/l of starch and 10 mg/l of hypochlorite and that when 1 ml is dosed into a litre of water sample, the dosage is 3 mg/l expressed in term of combination of hypochlorite and starch (i.e. in terms of polyelectrolytes itself).

115 Table 3: Expression of Polyelectrolyte (ml) in terms of Dosage (mg/l)

	Polyelectrolyte expressed as				
Volume of coagulant per litre					
	Starch (mg/l)	Hypochlorite (mg/l)	Polyelectrolyte (mg/l)		
(ml)					
0	0	0	0		
2.5	7.5	25.0	32.5		
5.0	15.0	50.0	65.0		
7.5	22.5	75.0	97.5		
10.0	30.0	100.0	130.0		
12.5	62.5	125.0	187.5		
15.0	75.0	150.0	225.0		

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117 Table 4 shows the relative performance of varying dosages of polyelectrolytes A, B and C under neutral 118 condition. The performance of polyelectrolyte A is generally better than those of B and C at neutral pH. 119 Polyelectrolyte B on the other hand, increases the turbidity of the water with which it was tested. The 120 percentage turbidity removals were computed based on Table 4 for polyelectrolytes A, B and C for each 121 experimental run. Table 5 shows that polyelectrolyte A at pH 7 performs generally better than 122 polyelectrolytes B and C. 123 124 125 126 127 128 129 130 131 132 Table 4: Relative effects of varying dosages of polyelectrolytes A, B and C on turbidity 133 removal and pH of treated water samples Polyelectrolyte C Polyelectrolyte B Polyelectrolyte A Dosage

(mg/l)	Final turbidity	Final pH	Final turbidity	Final pH	Final turbidity	Final pH
	(NTU)		(NTU)		(NTU)	
0						
15	13.2	7.8	42.2	7.6	6.2	7.7
32.5	11.6	7.7	43.4	7.6	4.8	7.7
65	9.8	7.6	44.0	7.5	3.8	7.5
97.5	10.0	7.4	44.0	7.4	4.2	7.4
130	12.4	7.3	44.7	7.4	5.6	7.4

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 Table 5:
 Summary of Percentage Turbidity Removal of Polyelectrolytes A, B and C

	Dosage (mg/l)	Percentage Turbidity Removal				
		Polyelectrolyte A	Polyelectrolyte B	Polyelectrolyte C		
	130	63.52	-18.26	45.48		
	97.5	70.39	-17.8	50.15		
	65	74.8	-16.44	52.68		
	32.5	70.48	-14.46	49.08		
	15	59.75	-12.95	43.84		
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141	3.2. Main Effect and Ir	nteraction of Chargin	ng Variables by Facto	orial Analysis		
142	The main and interaction	effect(s) were used to	determine the level of ir	fluence the factors had on one		

143 another and how the influence was occurring (Uduman et al., 2010). Table 6 shows the summary of this

results. It can be seen that power had the most significant main effect on the coagulation characteristics of the starch-based polyelectrolyte while the highest level of interaction effect was observed between the separating distance and the time of charging.

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_ 1	40	

Table 5: Summary of main effect and interaction effect of the charging variables

Effect	Polyelectrolyte A	Polyelectrolyte B	Polyelectrolyte C
Mean	74.75	27.98	52.63
Р	19.0	-21.07	17.25
D	-0.5	8.43	1.25
PD	0	2.17	-1.75
Т	1.0	5.97	1.25
PT	-3.5	-3.68	4.25
DT	-7.0	1.98	-8.75
PDT	2.5	-12.03	-0.75

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151 **4. CONCLUSION**

Based on the results, the following conclusions could be drawn: when 3 g of cassava starch and 10 g of calcium hypochlorite are dispersed in hot distilled water and charged, the resulting product, starch-based polyelectrolyte, is effective in coagulating raw water. A dosage of 65 mg/l of polyelectrolytes A and C result in 88% and 69% turbidity removal respectively. The same dosage of polyelectrolyte results in 7.5% turbidity addition.

Polyelectrolyte A performs generally as well as alum but does not depress the pH as alum does. This means that the starch-based polyelectrolyte needs little or no lime to aid it in the reduction of potable water at the neutral pH range. The solution is stable at room temperature. It neither spoils nor loses its coagulation effectiveness for at least 12 days. Power supplied during the charging process has the most significant effect on the coagulation characteristics of the starch-based polyelectrolyte while the highest level of interaction effect is between separating distance and time of charging.

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