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# Removal of Trimethoprim from Water using Carbonized Wood Waste as Adsorbents

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#### Abstract

Daniellia—*oliveri* sawdust-based adsorbents were employed to remove trimethoprim (TMP) from water. The sawdust was thermally carbonized and activated in-stu with ZnCl2 and  $H_3PO_4$  separately. The adsorbents surface features were profiled using scanning electron microscopic (SEM) and pH point of zero charge (pH<sub>*pzc*</sub>) analyses. The prospects of the adsorbents for the removal of trimethoprim from water were verified. The adsorption processes were performed under different experimental conditions. The adsorption isotherm, the kinetics, and the thermodynamics were studied; and the data fitting output revealed that both chemisorptions and physisorption occurred. Surface and pore diffusion played active role in the adsorption of TMP by the adsorbents. The optimum conditions for adsorption of TMP by the adsorbents were pH at slightly acidic to neutral medium and temperature at room temperature. The fitting isotherm models were: Langmuir ( $R^2 = 0.993$ ) for the zinc-chloride-activatedcarbon, Temkin ( $R^2 = 0.962$ ) for the phosphoric-acid-activated-carbon, and the kinetics: pseudo-second order ( $R^2 = 0.997$ ) for both. The maximum monolayer adsorption capacities of the adsorbents for TMP was 4.115 and 6.495 mg/g, respectively. The thermodynamic parameters determined suggested feasibility, spontaneity, and endothermicity of the adsorption processes. The results reveal that the adsorbents were good prospects for the removal of TMP from water.

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#### 1. Introduction

Necessities for procurement of means of livelihood, maintenance of quality life and attainment of luxurious status had prompted man to recreate the environment. Advancements in various frontiers of human endeavors had created lots of pollutants and contaminants.

Sources of pollutants and contaminants in the environment include: agricultural practices, medicine/pharmaceutical indus-

tries, automobile/mechanical works, transportation, construction, manufacturing, tourism, research, productions and so on [1]. The propensity of these hazardous substances into the lithosphere and hydrosphere was high as a result of their proximity to man. Most of these anthropogenic toxic materials upset the delicate balance of the ecosystem. The resultant of this upsetting was untoward environmental episodes.

The advancement of man in various facets has led to new products, generation of new by-products and new wastes. Some of new wastes had been identified as emerging contaminants of concerns (ECCs). The ECCs were of different categories:

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nano-materials [1] pesticides, herbicides, personal care products and pharmaceuticals (PCPPs) [2]. The pharmaceuticals are designed to fight specific ailment and/or microorganisms that cause diseases. When administered through involuntary exposure, the pharmaceuticals had been appraised to induce an array of health and ecological complications [3]. Some of the identified ecological complications include and not limited to: hermaphroditic feature in fish; regeneration impairment in injured toads; and drug resistance in microbe populations [4, 5, 6]. The environmental levels of pharmaceuticals are generally low to cause acute health effect on human. However considering chronic exposure, precautionary principle approach should be adopted to forestall incidence of health complications. The involuntary presence of pharmaceuticals should be prevented in public resources like drinking water and air to protect the vulnerable groups like children and populations with compromised immune system from involuntary exposure.

Pharmaceuticals were of different categories classified by their functions. One of the main categories with high dispensability was antibiotics e.g. trimethoprim (TMP) [7, 8, 9]. High dispensability of antibiotics was responsible for their high levels in the environment. Presence of antibiotics in the environment was highly undesirable as they induce drug resistance in microorganisms and this is a major factor of pandemic [10, 11]. TMP is an antibacterial and anti-malarial drug. It has reported half-lives in the range 5 - 100 days. It is insignificantly photodegraded, which suggests a relative persistence [12] in the environment. TMP is mutagenic, teratogenic, embryotoxic and folate antagonism [13]. Hence TMP should be removed from water resources.

The researchers are working to develop technologies [14] and strategies [15, 16, 17, 18] to remove various forms of pollutants and contaminants, including pharmaceuticals from water. Recently the most sustainable technology so far developed was adsorption process [19, 20, 21, 22, 23, 24, 25]. The trending strategy was the use of agro-wastes as adsorbents for the removal of pollutants and contaminants. In Nigeria, wood waste like sawdust remains one of the high volumes, posing great challenge to manage. In 2010, Nigeria generated over 1,000,000  $m^3$  (353,146.667 tonnes) [26] of sawdust and most managed by open air combustion. This combustion method led to generation of COx, NOx, SOx: radiation forcing substances [26]. Conversion of the sawdust to highly demanded product like adsorbent is both economically and environmentally sustainable strategy.

In this work, Daniellia-*oliveri* sawdust, a high volume agroprocessing waste in Nigeria and not yet reported for any usage, was processed and used to remove from contaminated water.

## 2. Materials and Methods

#### 2.1. Materials

Some of the materials used include *Daniellia-oliveri* sawdust (collected from a local factory, washed, dried and segregated), analytical grade ZnCl<sub>2</sub>,  $H_3PO_4$ , (purchased from local chemical dealer) TMP (primary standard, >98.5% purity) (supplied Bond Chemical Industries Limited, Awe, Oyo) and HCl. The equipment made used of were furnace (Carbolite AAF 1100), ultraviolet-visible (UV) (B-UV 1800PC) spectrophotometer, pH meter (Jenway 3520), and scanning electron microscope (SEM) (Aspex 3020) and oven (Carbolite).

## 2.2. TMP

The structural formula and the properties of trimethoprim are given in Figure 1 [27] and Table 1 [28] below.



Figure 1: Trimethoprim

#### 2.3. Preparation of Adsorbents

The *Daniellia*—*oliveri* sawdust was washed with plenty of distilled water to remove surface impurities and then sundry. The sample was dried in an oven at 70 °C for 72 hours; then ground and stored as Raw Daniellia Sawdust (RDS). The method was described elsewhere [29], 3g of RDS was mixed with 3 mL of 1 M H<sub>3</sub>PO<sub>4</sub> and 3 mL of 1 M ZnCl<sub>2</sub> each in separate crucibles. These were subjected to the furnace at 800 °C (carbonization and activation in self-generated atmosphere) for 5 minutes. The adsorbents were stored in air tight container as H<sub>3</sub>PO<sub>4</sub> activated carbon (PACB) and ZnCl<sub>2</sub> activated carbon (ZAC) respectively.

#### 2.4. Characterization of Adsorbents

SEM and  $pH_{pzc}$  analyses were carried out using methods [30] and [31] respectively to study the surface morphology and pH of point of zero charge of the adsorbents.

## 2.5. Batch Adsorption Studies

The triplicate optimized batch adsorption processes were carried out. The adsorption bottles containing adsorbent-adsorbate mixtures were shake on a mechanical shaker operating at 160 rpm. The effects of contact times (0 - 240 min), adsorbent dosages (0.05 - 0.5 g), pHs (2, 5, 8), initial adsorbate concentrations (15 - 50 mg/L), and temperatures (29 - 60 °C) were determined. After the processes reached equilibrium, the mixtures were filtered using filter papers and the filtrates analyzed at 282

Table 1: Properties of Trimethoprim

Common Names: IUPAC Names	Trimethoprim, Trimethoprime, trimethoprimum, trimetoprima, trimpex 5-[(3,45-trimethoxyphenyl)methyl]-2,4-pyrimidinediamine				
Molecular formula	$C_{14}H_{18}N_4O_3$				
Molecular weight	290.32 g/mol				
UN Number	2811				
Max $\lambda$	282 nm				
Solubility	400 mg/L (25°C)				
Toxicity	Vomiting, headache, mental depression, confusion, bone marrow depression				
рКа	7.12				
Description	Antibacterial, antimalaria				
Interaction sites	$\pi - \pi$ , $n - \pi$ , -NH <sub>2</sub> , $\equiv$ N:				

nm with the UV spectrometer to determine the absorbances of trimethoprim in solutions after adsorption. The absorbances were converted to corresponding concentrations using standard curve equation. Each experiment was performed in triplicate. Equations (1) and (2) were used to determine amount of adsorbate adsorbed per unit mass of adsorbent and percentage adsorbate removal respectively:

$$q_e = \frac{(C_i - C_e)V}{m} \tag{1}$$

% adsorbate removal = 
$$\frac{(C_i - C_e)V}{C_i} \times 100$$
 (2)

where  $q_e$ ,  $C_i$ ,  $C_e$ , m and V are the amount of trimethoprim adsorbed by the adsorbent at equilibrium (mg/g); the initial trimethoprim concentration (mg/L); the trimethoprim concentration at equilibrium (mg/L); the mass of the adsorbent (g); and the volume of the solution (L) respectively.

#### 2.6. Adsorption Isotherms

Table 2 contains selected isotherm models used to process experimental data and then described the adsorption processes.

The Langmuir Model: A plot of  $C_e/q_e$  versus  $C_e$  gives a straight line with intercept:  $1/K \cdot q_0$ ; and slope:  $1/q_0$ . Langmuir isotherm equilibrium parameter,  $R_L$ :  $R_L = \frac{1}{1+K_LC_i}$  [39].  $C_i$  = highest initial adsorbate concentration (mg/L);  $R_L > 1 =$  unfavourable;  $R_L = 1 =$  linear; favorable  $0 < R_L < 1 =$  favourable and  $R_L = 0 =$  irreversible. The Freundlich Model: A log-log plot should yield an intercept of log  $K_F$  and a slope of 1/n. n defines isotherm shape; 1/n = 0 = irreversible; (0 < 1/n < 1) = favourable; unfavourable (1/n > 1); n = 1 = linear; n > 1 = physical process and n < 1 = chemical process. The Temkin Model:  $K_T =$  Temkin isotherm equilibrium binding constant (L/g);  $b_T =$ 

Table 2: Isotherm Equat	tions
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Model	Linearised Formulae	Reference
Halsey	$\ln q_e = \frac{1}{n} \ln K - \frac{1}{n} \ln C_e$	[32]
Harkin-Jura	$\frac{1}{q^2} = \frac{B}{A} - \frac{1}{A} \log C_e$ $\frac{C_e}{q_e} = \frac{1}{K \cdot q_0} + \frac{C_e}{q_0}$	[33]
Langmuir		[34]
Freundlich	$\ln q_e = \ln K_F + \frac{1}{n} \ln C_e$	[35]
Temkin	$q_e = B \ln K_T + B \ln C_e$	[36]
Redlich-Peterson	$\log \frac{C_e}{q_e} = \log K_R +$	[37]
	$\beta_R T \log C_e$	
Dubinin–	$\ln q_e = \ln(q_s) - (K_{ad}\varepsilon^2)$	[38]
Radushkevich		

Temkin isotherm constant; R= universal gas constant (8.314 J/mol/K); T= Temperature at 298 K; B = Constant related to heat of sorption (J/mol). The Redlich-Peterson isotherm:  $C_e$  = equilibrium concentration;  $q_e$  = adsorption capacity of the adsorbent;  $\beta$  = desorption constant and  $K_R$  = R-P isotherm constant (g/L). The Dubinin-Radushkevich Model:  $\varepsilon = RT \ln[1 + \frac{1}{C_e}]$ ;  $E = [\frac{1}{\sqrt{2}B_{DR}}]$ ;  $K_{ad} = B_{DR}$ ; E<8 kJ/mol (physical adsorption); E = 20–40 kJ/mol (chemical adsorption) [38]; E = 8–16 kJ/mol (ion-exchange) [40];  $q_e$  = amount of adsorbate on the adsorbent at equilibrium(mg/g);  $q_s$  = theoretical isotherm saturation capacity (mg/g);  $K_{ad}$  = Dubinin-Radushkevich isotherm constant (mol<sup>2</sup>/kJ<sup>2</sup>).

### 2.7. The Kinetics

Samples were taken from adsorption bottles at time intervals and the amounts of adsorbates were measured. The amount of adsorbed adsorbate at time t,  $q_t$  (mg/g), was calculated using Equation 3 and kinetic parameters were calculated using kinetic equations presented in Table 3.

$$q_t = \frac{(C_0 - C_t) V}{m} \tag{3}$$

The Pseudo-First Order:  $q_e$  and  $q_t$  are the amounts of the dye adsorbed (mg/g) at equilibrium and at time t (min), respectively, and  $k_1$  is the rate constant adsorption (min<sup>-1</sup>). A plot of  $\log(q_e - q_t)$  versus t gives a slope of  $\left(-\frac{k_1}{2.303}\right)$  and intercept of  $\log q_e$ . The Pseudo-Second Order:  $h = k_2 \cdot q_e^2$ ;  $q_e$  is the amount of the solute adsorbed at equilibrium per unit mass of adsorbent (mg/g),  $q_t$  is the amount of solute adsorbed (mg/g) at any given time, t (min) and  $k_2$  is the rate constant for pseudosecond-order adsorption (g/mg/min) and h, known as the initial sorption rate. The values of  $q_e$  and  $k_2$  can be obtained from the slope and intercept of the plot of  $t/q_t$  versus t respectively. If the sorption follows pseudo-second order, h, is described as the initial rate constant as t approaches zero. The Elovich:  $\alpha$  is the initial adsorption rate (mg/g. min),  $\beta$  is the desorption constant (g/mg) and  $q_t$  is the amount of solute adsorbed (mg/g) at any given time, t (min). A plot of  $q_t$  versus ln t gives a straight line with intercept  $\frac{1}{\beta} \ln(\alpha\beta)$  and slope  $\left(\frac{1}{\beta}\right)$ . Intra-Particle Diffusion:  $q_t (mg/g) = quantity of adsorbate adsorbed at time, t and$  $k_{id}$  (mg/gh<sup>0.5</sup>) = intra-particle diffusion constant. A graph of  $q_t$ versus  $t^{0.5}$  gives a straight line with slope,  $k_{id}$ , and intercept, x. x defines the thickness of the boundary layer. The Boyd Kinetic: B =  $\frac{\pi^2 D_i}{r^2}$ ; B is a constant; D<sub>i</sub> (m<sup>2</sup>/s) is diffusion coefficient; r is radius of adsorbent particle; F is the fraction of adsorbate adsorbed at given time, t, and Bt is a function of F. If graph of  $[-0.4977 - \ln(1 - F)]$  vs time, t, is linear and intercept = 0, then the determining step of the adsorption process is intraparticle diffusion; but if intercept  $\neq 0$ , the determining step may be film diffusion.

#### 2.8. Thermodynamic Study

Experiments were performed at 29, 40, 50, 60 and 70  $^{\circ}$ C to establish the effect of temperature on the adsorption capacities of ZAC and PACB for the trimethoprim. The thermodynamic

Table 3: Kinetic Equations

Kinetic	Equation	Reference
Pseudo-First Order	$\log(q_e - q_t) = \log q_e - \frac{k_1}{2.303} .t$	[41]
Pseudo- Second Order	$\frac{t}{q_t} = \frac{1}{h} + \frac{t}{q_e}$	[42]
Elovich	$q_t = \frac{1}{\beta} \ln(\alpha\beta) + \left(\frac{1}{\beta}\right) \ln t$	[43, 44]
Intra-Particle Diffusion	$q_t = k_{id}t^{0.5} + x$	[45]
Boyd	$-0.4977 - \ln(1 - F) = Bt$	[46]

$$\Delta G^0 = -2.303RT \log K \tag{4}$$

$$\log\left(\frac{C_{Ae}}{C_e}\right) = \frac{\Delta S^0}{2.303R} - \frac{\Delta H^0}{2.303RT}$$
(5)

 $K = \left(\frac{C_{Ae}}{C_e}\right); C_{Ae} = (C_i - C_e) (mg/L) = amount of trimethoprim$  $adsorbed on adsorbent at equilibrium; <math>C_e (mg/L) = amount$ of trimethoprim in solution at equilibrium; R = gas constant (8.314 J/mol/K); T = temperature (K). The enthalpy change ( $\Delta$ H), the entropy change ( $\Delta$ H) and change in saturd free energy ( $\Delta$ G) were calculated. The spontaneity of the processes was thereby determined.

#### 3. Results and Discussion

#### 3.1. Morphology of ZAC and PACB

Figure 2 showed characteristic surface morphology of the adsorbents. Carbonization and activation processes impacted the adsorbents' surfaces with characteristic morphology and chemical profile. Using Gwyddion 2.23-1 application for SEM image processing, the average pore diameters respectively of ZAC and PACB were approximately 30 and 55 nm. High temperature of carbonization removed volatile compounds in the matrix of raw sawdust thereby created pores. High temperature could also break down cellulosic contents of the precursor sawdust and aided aromatization of carbon chains. High temperature chemical activation effected dehydration and decarboxylation of sawdust. It could be deduced from the changes in the surface morphology after adsorption that pores were active in adsorbing trimethoprim molecules.

## 3.2. Determination of Equilibrium Time and Effect of Contact Time on Trimethoprim Adsorption

The amount of TMP adsorbed per unit mass (q) of each of ZAC and PACB increased rapidly within first 40 min. The q thereafter increased slowly and insignificantly over next two hours for both systems. After 60 min, almost all the available adsorption sites had been occupied for both systems. The qat time t ( $q_t$ ) for the TMP-ZAC system was 1.123 mg/g within first 60 min while it was 0.272 mg/g for the 180 min afterwards. The  $q_e$  for the TMP-PACB system was 0.996 mg/g in first 60 min while 0.204 mg/g for the 180 min after first 60 min were economically insignificant. The economic time of adsorption for both systems was 60 min.

## 3.3. Effect of Adsorbent Dose on Trimethiprim (TMP) Adsorption

The increase in mass of each of ZAC and PACB led to decreased amount of TMP adsorbed per unit mass during adsorption processes. The increased mass of the adsorbents provided more sites of adsorption for constant amount of TMP molecules. ZAC and PACB adsorbed equal amounts of the



Figure 2: (a) PACB (before adsorption); (b) PACB (after adsorption); (c) ZAC (before adsorption); (d) ZAC (after adsorption) at magnification × 500.

TMP per unit mass at 0.15g each (at the point of intersection) (Figure 3b). 0.2, 0.3 and 0.4 g of ZAC adsorbed 68 %, 82 % and 87 %; and PACB adsorbed 65 %, 76% and 75% of TMP respectively at equilibrium time. The equilibrium dose was reached at 0.3 g for both adsorbents. The economic dose for ZAC was 0.3 g while 0.2 for PACB.

#### 3.4. Effect of pH on Trimethoprim Adsorption

The  $pH_{pzc}$  is the pH of solution at which electrical charge of a solid surface in the solution is neutral. The pKa of trimethoprim and the  $pH_{pzc}$  of the adsorbents are parameters whose electrical charge changes with pH of solution. The maximum attraction between the adsorbate and the adsorbents would take place when electrical charges on them are opposite. TMP is a basic compound with pKa 7.12. At pH above 7.12, TMP was neutral while at pH below 7.12 TMP was positively charged (the hypothesized interactions were presented in Equations 6 to 10). At pH below 7.12, the oxygen and nitrogen atoms on TMP donated lone pairs of electrons to the protons in solution and TMP became positively charged. At pH above 7.12, the lone pairs of electrons on TMP oxygen and nitrogen atoms repelled lone pairs of electrons on hydroxyl groups in solution and TMP remained neutral. At about pH 7, TMP was at the threshold of changing from positively charged (+) to neutral (0), ZAC (pH<sub>pzc</sub> 7.6) positively charged (+) while the PACB

(pH<sub>*pzc*</sub> 6.6) negatively charged (-). At pH below 6, the adsorbents and trimethoprim were all positively charged and the repulsive effect de-enhanced adsorption. At pH above 8, lone pairs of electrons on neutral TMP repelled negatively charged adsorbents' surface and chemisorption was de-enhanced. The maximum interactions occurred at slightly acidic to neutral pH (pH 6) where electrical charges on the adsorbents and trimethoprim were opposite (Figure 3c). At pH 2.5, ZAC adsorbed 1.262 mg/g of TMP; at pH 5, 1.266 mg/g and at pH 8, 1.25 mg/g. And for PACB, at pH 2.5, 1.164 mg/g of TMP was adsorbed, at pH 5, 1.215 mg/g and at pH 8, 1.138 mg/g.

$$TMP (pKa) \xrightarrow{pH>7.12} TMP(O)$$
(6)

$$Me\ddot{O} \equiv N^{\cdot}(TMP) + OH^{-} \to Me\ddot{O}$$
(7)  
$$\equiv N^{\cdot}(TMP) \leftrightarrow OH^{-} (neutralTMP)$$

$$TMP (pKa) \xrightarrow{pH<7.12} TMP(+)$$
(8)

$$MeO \equiv N^{:}(TMP) + 2H^{+} \rightarrow MeOH^{+}$$
(9)  
$$\equiv NH^{+} (positively charged TMP)$$

## 3.5. Effect of Temperature on Trimethoprim Adsorption

TMP is a biphenyl compound with functional group branches. The bulky structure of TMP may constitute stearic hindrance to

pore accessibility. At elevated temperatures, TMP molecules acquired more kinetic energy which led to frequent knocking with the adsorbents' pore openings. The impact of the knocking might 'mend and tend' the molecules or wear the pore openings and eventually forced the molecules into the pores. The higher the temperature, the higher was the kinetic energy and the contact between molecules and pores. At equilibrium, 1.6 mg/g (64 %) and 1.904 mg/g (76 %) of TMP were adsorbed by ZAC at 29 °C (room temperature) and 60 °C respectively. Also 1.196 mg/g (72 %) and 1.326 mg/g (80 %) adsorbed respectively at 29 °C and 60 °C by the PACB (Figure 3d). The enhanced adsorption of TMP on ZAC and PACB as a result of increase in temperature was economically unfavourable. From economic perspective, room temperature would be chosen considering costs of heating and cooling the system after adsorption before discharge, or environmental impact of discharged warm/hot wastewater. It would be less expensive to subject the TMP treated water to secondary adsorption process.

#### 3.6. Effect of initial TMP Concentrations

The  $q_e$  (mg/g) of the adsorption of TMP onto ZAC and PACB increased as the initial TMP concentrations increased. This was due to the more TMP molecules in the solutions which were in contact with constant active adsorption sites on the adsorbents. For the TMP-ZAC system, at 15, 20, 25, 30, 35, 40 and 50 mg/L, 82.6 %, 83.5 %, 82.8 %, 81 %, 79.5 % 75 % and 70 % of TMP were adsorbed respectively. And for the TMP-PACB system, at 15, 20, 25, 30, 35, 40 and 50 mg/L, 75 %, 77.9 %, 81.7 %, 79.9 %, 78.8 %, 80 % and 75 % were adsorbed respectively (Figure 3e). A unit mass of ZAC had highest removal efficiency for TMP at 20 mg/L of solution while PACB at 25 mg/L of solution.

#### 3.7. Adsorption Isotherms for TMP Adsorption

The isotherms applied to describe the adsorption of TMP onto ZAC and PACB were Langmuir, Freudlich, Temkin, Redlich-Peterson, Dubinin-Radushkevich, Harkin-Jura and Halsey. The isotherms that best fit were Langmuir (R<sup>2</sup>=0.993) and Temkin  $(R^2=0.962)$  for TMP-ZAC adsorption and TMP-PACB respectively (Table 4). These suggested that TMP-ZAC adsorption system was monolayer and TMP-PACB multilayer. The Langmuir isotherm equilibrium parameter, R<sub>L</sub> values: 0.108 and 0.206 for the TMP-ZAC and the TMP-PACB meant that both processes were favorable. The adsorption affinity  $(K_L)$  for ZAC-TMP process was higher, which implied that it was more favorable compare to PACB-TMP process. The  $q_{max}$  for ZAC was 4.115 mg/g and PACB 6.495 mg/g (Table 6). The multilayer coverage exhibited by the TMP-PACB process explained its higher q<sub>max</sub>. Also wider pores exhibited by PACB (Figure 2) might be more accessible to TMP molecules than for ZAC. Comparison of qmax of some adsorbents reported in literatures indicated that ZAC and PACB were better adsorbents for TMP adsorption from aqueous solution. The 1/n as determined from the Freundlich Isotherm for the TMP-ZAC and the TMP-PACB processes were 0.472 g/L and 0.636 g/L respectively. These further confirmed the processes to be favorable. Both processes were physical because n greater than 1 for

Table 4: Isotherm parameters for TMP adsorption onto the ZAC and the PACB

Isotherm	ZAC	PACB
Langmuir		
$q_0 (mg/g)$	4.115	6.495
$k_L (L/mg)$	0.165	0.077
$R_L$	0.108	0.206
$R^{\overline{2}}$	0.993	0.913
Freundlich		
$K_F (mg/g)$	0.849	0.653
n (L/g)	2.120	1.572
1/n	0.472	0.636
$\mathbb{R}^2$	0.959	0.955
Temkin		
$K_T (mg/L)$	1.306	1.673
$b_T$ (KJ/mol/g.L)	2.531	0.664
β (L/g)	0.992	1.500
$R^2$	0.988	0.962
Redlich-Peterson		
$K_R$ (g/L)	1.178	1.528
$\beta_R$ (g/mg.K)	0.0017	0.0013
$R^2$	0.966	0.879
Dubinin-Radushkevich		
$q_s (mg/g)$	2.821	3.320
$K_{ad}$ (JL/mol.mg)	$2 \times 10^{-6}$	$1 \times 10^{-8}$
E (KJL/mol.mg)	0.50	2.5
R <sup>2</sup>	0.966	0.921
Harkin-Jura		
$q_e (mg/g)$	1.287	1.220
$A (mg^3/g^3)$	1.783	1.745
R <sup>2</sup>	0.840	0.908
Halsey		
n (g/L)	-2.105	-1.572
K (mg/L)	1.415	0.668
$R^2$	0.959	0.957



Figure 3: Effect of (a) contact time, (b) adsorbent dose, (c) solution pH, (d) temperature and (e) initial adsorbate concentration on the adsorption of TMP onto ZAC and PACB

each (Table 4). Adsorption intensity (n) was higher for ZAC-TMP process which in agreement with explanation given under Langmuir isotherm. 1/n, a function of surface heterogeneity, was higher for PACB-TMP system. Positive values of bonding energy ( $b_T$ ) obtained from Temkin isotherm for both processes indicated endothermic processes. The adsorption energy, E (kJ/mol), derived from Dubinin-Radushkevich isotherm for the TMP-ZAC and the TMP-PACB processes were 0.50 and 2.5 respectively. These values described both processes as physical. Halsey isotherm described a heterogeneous adsorbent surface and multilayer adsorption system. Relatively close values of adsorption coefficients of both Halsey and Temkin isotherms further established PACB-TMP system as multilayer adsorption and PACB had heterogeneous surface. As could be deduced from the coefficient values of the isotherm and kinetic (section 3.9 below) models, the adsorption of TMP by ZAC and PACB involved both chemisorption and physisorption. Chemisorption involves reaction among functional groups on both the adsorbents (ZAC and PACB) and the adsorbates (TMP). Hydroxyl group would like form hydrogen bond with adsorbents' N and



Figure 4: van't Hoff plot of  $\log K$  vs 1/T adsorption of TMP onto (a) ZAC and (b) PACB.

Table 6: Comparison of maximum monolayer adsorption capacities of different adsorbents for TMP

Adsorb	ent Adsorbent	capacity Reference
	(mg/g)	
GAC	0.51	[47]
PAC	0.51	[47]
Clay	47	[48]
BHB	0.508	[49]
LZB	0.294	[49]
ZAC	4.115	This work
PACB	6.495	This work
CAC	amonulan active	tad aamhan

GAC – granular activated carbon PAC – powdered activated carbon sediment BHB; sediment LZB

O bearing functional groups, and vice versa:

$$NH_2 - TMP - NH_2 + HO - ZAC/PACB \rightarrow H_2N - TMP$$
 (10)  
-  $HN \cdots HO - ZAC/PACB$ 

Multilayer adsorption of TMP onto PACB was as a result of reaction between electron withdrawing and electron donating functional groups on already adsorbed TMP and free adsorbate molecules in solution:

$$H_2N - TMP - HN \cdots HO - PACB + H_2N - TMP \rightarrow (11)$$
  
$$\rightarrow TMP - H_2N \cdots H_2N - TMP - H_2N \cdots OH - PACB$$

 $\pi$ - $\pi$  and n- $\pi$  are electron density centers capable of facilitating electrostatic interactions between TMP molecules and the adsorbents through resonance of conjugated double bonds and aromaticity of benzene rings:

$$TMP - N^{-}-C = N^{+}H_{2} + O^{-}C_{6}H_{6} - PACB/ZAC \rightarrow (12)$$
$$TMP - N^{+} \cdots O^{-}C_{6}H_{6} - PACB/ZAC$$

## 3.8. Kinetic Studies of Adsorption of TMP onto ZAC and PACB

The data of adsorption of TMP on ZAC and PACB were tested with0 five different kinetic models: Pseudo-first order,

Table 5: Kinetic parameters for TMP adsorption onto the ZAC and the PACB

Kinetic	ZAC	PACB
$q_e, e_{xp} (mg/g)$	1.395	1.200
Pseudo-first order		
$q_e (mg/g)$	0.670	0.581
$K1 (min^{-1})$	0.014	0.014
$\mathbb{R}^2$	0.964	0.920
Pseudo-second order		
$q_e (mg/g)$	1.425	1.230
$h(min^{-1})$	0.107	0.097
K2	0.053	0.064
$\mathbb{R}^2$	0.997	0.997
Elovich		
A	0.535	0.445
$\beta$ (min.g/mg)	4.40	5.05
$\mathbb{R}^2$	0.939	0.969
Intraparticle diffusion		
x (mg/g)	0.594	0.518
$K_{id}(mg/g.min^{1/2})$	0.061	0.053
$\mathbb{R}^2$	0.877	0.888
Boyd		
X	1.475	1.617
B (min.g/mg)	-0.004	0.003
R <sup>2</sup>	0.211	0.017

	$\Delta G^0 (kJ/mol/K)$			$\Delta S^0 (J/mol/K)$	$\Delta H^0 (kJ/mol/K)$	
	302	313	323	333		
ZAC	-1.444	-2.39	-2.64	-3.22	+54.60	+14.99
PACB	-2.342	-2.74	-2.90	-3.76	+42.16	+10.46

Pseudo-second order, Elovich, Intraparticle and Boyd. Both TMP-ZAC and the TMP-PACB processes data fitted pseudosecond order equation with  $R^2 = 0.997$  (Table 5). The rate of adsorption in both processes depended on the states of both the adsorbate and the adsorbents. The initial rate constants for ZAC and PACB, h (mg/g/min), were 0.107 and 0.097 respectively, implied that rate at which TMP adsorbed on to the ZAC was higher initially. However, overall rate constant of PACB-TMP process was higher. High adsorption coefficient  $(R^2)$  for Elovich model suggested a level of chemisorptions for both systems. Higher Elovich  $R^2$  for PACB-TMP process also indicated considerable multilayer coverage compare to ZAC-TMP process. However, earlier it was explained from the isotherm data fitting that adsorption of TMP by the adsorbents were both chemisorptions and physisorption. High coefficient values for pseudo first order kinetic and relatively high R<sup>2</sup> for intraparticle diffusion model were pointers to physical adsorption. Surface and pore diffusions played important role in the adsorption of TMP by ZAC and PACB. PACB having wider average pore diameter than ZAC could easily accommodate bulky TMP molecules and hence higher qmax. Diffusion of TMP molecules at the surface of the adsorbents occurred rapidly while diffusion down the adsorbents' pores took place steadily until equilibrium is attained.

#### 3.9. Thermodynamic Study of the TMP Adsorption

The functions of temperature, the standard enthalpy ( $\Delta H^0$ ), the standard entropy ( $\Delta S^0$ ) and the standard Gibb's free energy ( $\Delta G^0$ ) of the adsorption processes were investigated for spontaneity and feasibility. The van't Hoff plot of graph of log K versus 1/T gave the slope and the intercept as  $\Delta H^0$  and  $\Delta S^0$  respectively (Table 7, Figure 4). The negative values for  $\Delta G^0$  at all investigated temperatures indicated that the adsorption of TMP onto ZAC and PACB were spontaneous. The positive value of  $\Delta S^0$  showed high degree of randomness of the processes while positive value of  $\Delta H^0$  indicated endothermic processes.

#### 4. Conclusion

This study verified the prospects of the processed Daniella oliveri sawdust as adsorbents for adsorption of TMP from aqueous solution. The economic time of adsorption of TMP by both adsorbents was 60 minutes while economic doses of adsorbents were 0.3 and 0.2 g for ZAC and PACB respectively at 29 °C. The  $q_{max}$  of both the ZAC and the PACB was obtained at slightly acidic-neutral pH. The ZAC data fitted best into the Langmuir isotherm and the PACB, the Temkin while data for both fitted best the pseudo-second order kinetic. Multilayer adsorption and average pore diameter were suspected to be responsible for higher  $q_{max}$  in PACB. The  $\Delta G^0$  values indicated that the adsorption processes were spontaneous and feasible. Also both reactions were endothermic and physical. The study revealed that Daniellia—*oliveri* sawdust based activated carbons were promising adsorbents for removal of TMP from water.

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#### References

- S. Hong, C. Ning, L. Jianguang, W. Dawei & X. Weixue "Preparation of Clay Nanocomposites Matrix", Journal of Chemical Science and Technology 2 (2013) 128.
- [2] Q. Sui, X. Cao, S. Lu, W. Zhao, Z. Qiu & G. Yu "Occurrence, sources and fate of pharmaceuticals and personal care products in the groundwater: A review", Emerging Contaminants 1 (2015) 14, https://doi.org/10.1016/j.emcon.2015.07.001.
- [3] Y. Luo, W. Guo, H. H. Ngo, L. D. Nghiem, F. I. Hai, J. Zhang & X. C. Wang "A review on the occurrence of micropollutants in the aquatic environment and their fate and removal during wastewater treatment", Science of the Total Environment 473 (2014) 619, https://doi.org/10.1016/j.scitotenv.2013.12.065.
- [4] D. C. Bean, D. M. Livermore, I. Papa & L. M. Hail "Resistance among Escherichia eoli to sulphonamides and other antimicrobials now little use in man", J. Antimicrobchemother 56 (2005) 962.
- [5] C. L. Ventola "The antibiotic resistance crisis: part 1: causes and threats", Pharmacy and Therapeutics 40 (2015) 277.
- [6] A. J. Ebele, M. A. E. Abdallah & S. Harrad "Pharmaceuticals and personal care products (PPCPs) in the freshwater aquatic environment", Emerging Contaminants (2017) 13, https://doi.org/10.1016/j.emcon.2016.12.004.
- [7] D. Maddileti, B. Swapna & A. Nangia "Tetramorphs of the antibiotic drug trimethoprim: characterization and stability", Cryst Growth Des 15 (2015) 1745, https://doi.org/10.1021/cg501772t.
- [8] F. J. Peng, G. G. Ying, Y. S. Liu, H. C. Su & L. Y. He "Joint antibacterial activity of soil-adsorbed antibiotics trimethoprim and sulfamethazine", Sci Total Environ 506 (2015) 58, DOI:10.1016/j.scitotenv.2014.10.117.
- [9] T. Thomas "Sulfamethoxazole/Trimethoprim ratio as a new marker in the raw wastewaters: a critical review", Science of the total Environment (2020), doi: 10.1016/j.scitotenv.2020.136916.
- [10] P. P. Gao, D. Q. Mao, Y. Luo, L. M. Wang, B. J. Xu & L. Xu "Occurrence of sulfonamide and tetracycline-resistant bacteria and resistance genes in aquaculture environment", Water Res 46 (2012) 2355, DOI: 10.1016/j.watres.2012.02.004.
- [11] E. Pradhan, S. Bhandari & R.E. Gilbert "Antibiotics versus no treatment for toxoplasma retinochoroiditis", Cochrane Database Syst Rev 5 (2016), doi: 10.1002/14651858.CD002218.pub2.
- [12] O. S. Jurg "An environmental risk assessment for human-use trimethoprim in European surface waters", Antibiotics 2 (2013) 115-162, Doi:10.3390/antibiotics2010115.

- [13] J. O. Straub "An Environmental Risk Assessment for Human-Use Trimethoprim in European Surface Waters", Antibiotics 2 (2013) 115– 162, doi: 10.3390/antibiotics2010115.
- [14] D. Dolar, N. Drašinac, K. Košutić, I. Škorić & D. Ašperger "Adsorption of hydrophilic and hydrophobic pharmaceuticals on RO/NF membranes: identification of interactions using FTIR", J. Appl Polym Sci 134, (2017), https://doi.org/10.1002/app.44426.
- [15] E. C. Salihi & M. Mahramanlioglu "Equilibrium and kinetic adsorption of drugs on bentonite: presence of surface active agents effect", Appl Clay Sci 101 (2014) 381, doi:10.1016/j.clay.2014.06.015.
- [16] C. Avila & J. García "Pharmaceuticals and Personal Care Products (PPCPs) in the Environment and Their Removal from Wastewater through Constructed Wetlands", Elsevier Science & Technology 195 (2015), https://doi.org/10.1016/B978-0-444-63299-9.00006-5.
- [17] Q. Liu, M. Li, F. Zhang, H. Yu, Q. Zhang & X. Liu "The removal of trimethoprim and sulfamethoxazole by a high infiltration rate artificial composite soil treatment system", Frontiers of Environmental Science and Engineering 11, 12 (2017), https://doi.org/10.1007/s11783-017-0920-z.
- [18] M. Samuel, M. Rosa, G. Jairo, S. Joanna, G. Sebastiano, R.C. Juan & P.O. Maria "Towards the removal of antibiotics detected in wastewaters in the POCTEFA territory: occurrence and TiO<sub>2</sub> photocatalytic pilot-scale plant performance", Water, **12** (2020) 1453, doi: 10.3390/w12051453.
- [19] J. T. Wang, J. Hu & S. W. Zhang "Studies on the sorption of tetracycline onto clays and marine sediment from seawater", J. Colloid Interface Sci 349 (2010) 578, https://doi.org/10.1016/j.jcis.2010.04.081.
- [20] Y. L. Zhang, S. S. Lin, C. M. Dai, L. Shi & X .F. Zhou "Sorptiondesorption and transport of trimethoprim and sulfonamide antibiotics in agricultural soil: effect of soil type, dissolved organic matter, and pH", Environ Sci Pollut Res 21 (2014) 5827, https://doi.org/10.1007/s11356-014-2493-8.
- [21] J. R. Li, Y. X. Wang, X. Wang, B. L. Yuan & M. L. Fu "Intercalation and adsorption of ciprofloxacin by layered chalcogenides and kinetics study", J. Colloid Interface Sci 453 (2015) 69, DOI: 10.1016/j.jcis.2015.03.067.
- [22] H. Liu, J. Zhang, H. H. Ngo, W. Guo, H. Wu, Z. Guo, C. Cheng & C. Zhang "Effect on physical and chemical characteristics of activated carbon on adsorption of trimethoprim: mechanisms study", RSC Adv 5 (2015) 85187, https://doi.org/10.1039/C5RA17968H.
- [23] J. Li & H. Zhang "Adsorption-desorption of oxytetracycline onmarine sediments: kinetics and influencing factors", Chemosphere 164 (2016) 156, https://doi.org/10.1016/j.chemosphere.2016.08.100.
- [24] M. Pan & L. M. Chu "Adsorption and degradation of five selected antibiotics in agricultural soil", Science of the Total Environment 545 (2016) 48, doi: 10.1016/j.scitotenv.2015.12.040.
- [25] M. Ming-sheng, M. Shuai-shuai, S. Li, K. Qiang & L. Yu-zhen "Adsorption characteristics of antibiotics trimethoprim by activated carbon developed from low-cost alligator weed: kinetics, equilibrium and thermodynamic analyses", Desalination and Water Treatment, (2017) 1, doi: 10.5004/dwt.2017.21053.
- [26] U. P. Onohie, E. K. Orhorhoro & P. E. Oyiboruona "Economic Potential and Benefits of Sawdust in Nigeria", International Journal of Research Publication 9 (2018) 1.
- [27] A. Reem, Al-Bayati, S. Athraa & Ahmed "Adsorption Desorption of Trimethoprim Antibiotic Drug from Aqueous Solution by Two Different Natural Occurring Adsorbents", International Journal of Chemistry 3 (2011), doi:10.5539/ijc.v3n3p21.
- [28] PubChem Trimethiprim. National Center for Biotechnology Information.

U.S. National Library of Medicine

- [29] A. A. Giwa, S. A. Adesokan & I. A. Bello "Adsorption of pyrimethamine from wastewater using activated carbons prepared from Daniellia-*oliveri* sawdust", Inter. J. Env. Anal. Chem 1, (2021), DOI:10.1080/03067319.2021.1884858.
- [30] R. Malik, D. S. Ramteke & S. R. Wate "Physico-Chemical and Surface Characterization of Adsorbent Prepared From Groundnut Shell by ZnCl<sub>2</sub> Activation and its Ability to Adsorb Colour", India journal of chemical technology (2006) 319.
- [31] R. M. Shrestha, A. P. Yadav, B. P. Pokharel & R. R. Pradhananga "Preparation and Characterization of Activated Carbon from Lapsi (*Choerospondias axillaris*) Seed Stone by Chemical Activation with Phosphoric acid", Res. J. Chem. Sci 2 (2012) 10.
- [32] G. Halsey "Physical Adsorption on non-uniform surfaces", J. Chem. Phys 16 (1948) 931-937.
- [33] W. D. Harkins & W. D. Jura "Adsorption equation", J. Chem, Phys 11 (1943) 430.
- [34] I. Langmuir "The adsorption of gases on plane surfaces of glass, mica and platinum", J. Am. Chem. Soc **40** (1918) 1361.
- [35] H. M. F. Freundlich "Over the Adsorption in Solution", J. Phys. Chem 57 (1906) 385.
- [36] M. I. Tempkin & V. Pyzhev "Kinetics of Ammonia Synthesis on Promoted Iron Catalyst", Acta Phys. Chim. USSR 12 (1940) 327.
- [37] C. H. Giles, D. Smith & A. A. Huitson "A general treatment and classification of the solute adsorption isotherm", Theoretical, J. Colloid Interface Sci 47 (1974) 755.
- [38] W. Rieman & H. Walton "International series of monographs in analytical chemistry, 38: Ion exchange in analytical chemistry", Oxford: Pergamon Press (1970).
- [39] W.T. Weber & K.P. Chakraborty "Pore and solid diffusion model for fixed bed adsorbent", J. Am. Inst. Chem. Eng 20 (1974) 228.
- [40] F. Helfferich "Ion exchange" New York: McGraw-Hill Book Co (1962).
- [41] S. Lagergren "Zur theorie der sogenannten adsorption gelöster stoffe", Kungliga Svenska Vetenskapsakademiens. Handlingar Band 24 (1898) 1.
- [42] Y. S. Ho & G. McKay, Can J Chem Eng 76 (1998) 822.
- [43] D. L. Sparks "Kinetics of Reactions in Pure and Mixed Systems in Soil Physical Chemistry", CRC Press, Florida (1986) 21.
- [44] F. E. Okiemen & V. U. Onyega "Binding of Cadmium, Copper, Lead and Nickel Ions with Melon (*Citrullus vulgaris*) Seed Husk", Biological waste 29 (1989) 11.
- [45] J. W. Weber & C. J. Morris "Proceedings of the International Conference on Water Pollution Symposium", Pergamon Press, Oxford 2 (1962) 231-266.
- [46] G. Boyd, A. Adamson & L. Myers "The exchange adsorption of ions from aqueous solutions by organic zeolites. II kinetics", J Ame. Chem. Soc 69 (1974) 2836.
- [47] A. O. Michael, O. M. Vincent, O. W. Shem, M. N. Holiness, W. M. Charles & U. Mamo "Adsorption Studies of Trimethoprim Antibiotic on Powdered and Granular Activated Carbon in Distilled and Natural Water", Int. J. S. Res. Sci. Engg. Technol 4 (2018) 223, DOI : https://doi.org/10.32628/IJSRSET1841117.
- [48] A. H. Al-Shukrawi, Y. A. Al-Baitai & H. D. Fadhel "Characteristics of Trimethoprim Adsorption on Attapulgite Iraqi Clay", J Chem Eng Process Technol 8 (2017), DOI: 10.4172/2157-7048.1000365.
- [49] L. Jia & Z. Hua "Factors influencing adsorption and desorption of trimethoprim on marine sediments: mechanisms and kinetics", Environ Sci Pollut Res 24 (2017) 21929. DOI 10.1007/s11356-017-9693-y.