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## Adsorption Kinetic of Mn(II) Ions in Synthetic Acid Mine Water Using Calcium Carbide Residue as an Adsorbents

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This investigation concentrated on diminishing the centralization of Mn(II) particles in manufactured corrosive mine water by the adsorption strategy utilizing calcium carbide buildups as an adsorbent. The adsorption procedure was examined utilizing 100 mL of arrangement of engineered corrosive mine with differing pH of 2.5 to 5 and weight of calcium carbide buildup from 2.5 to 7.5 g. The blend was mixed utilizing a shaker with speed of 200 rpm for 180 minutes with contact time ranges (5, 10, 20, 30, 40, 50, 60, 90, 120, 150, and 180 minutes) and the underlying centralizations of Mn(II) of between 20 to 100 mg/L. The outcomes demonstrated that adsorption limit of adsorbent increments with expanding of starting grouping of Mn(II) particles in the arrangement. The Freundlich isotherm model matches the observational information. Adsorption energy demonstrated that this response pursues the pseudo-second adsorption model. The most extreme adsorption limit with respect to Mn(II) as arrangement is 41.67 mg/g at pH 5. The outcomes additionally shown that calcium carbide buildups are reasonable for adsorbing Mn(II) particles from fluid arrangements.

**Keywords:** Adsorption, Kinetic, Calcium Carbide Residue, Mn Metal.

### 1. INTRODUCTION

The mining business is commonly a noteworthy supporter of corrosive mine seepage (AMD). Corrosive mine water that requirements genuine consideration when released into water bodies. One of the broke down metals in corrosive mine water (AMD) is manganese ions (Mn(II)) which causes dark brown shade of the water when reaching with air [1], and will be dangerous when the centralization of Mn(II) particles disintegrated in water more prominent than 0.02 mg/L [2]. Manganese metals particle in corrosive mine water are found in different structures, both intricate and natural mixes [3]. Manganese particles are hard to expel from AMD since they precipitated when the pH is over 10 [4]. Accordingly, it is a need to diminish the manganese particles so the water is ok for condition. A few innovations have been created for killing corrosive mine water, including the adsorption procedure, particle trade, invert assimilation dissolvable extraction, flocculation and film detachment [5], electrochemistry [6], electrodialysis [7], just as ultrafiltration innovation, electrolysis, and extraction [8]. Among these techniques, adsorption is financially and in fact straight forward, moderately simple to work, basic structure, does not cause

unsafe and exceptionally powerful in adsorbing substantial metals particles [9, 10].

Enacted carbon is a standout amongst the best adsorbents, yet it requires costly expenses in the actuation procedure. In this way, it is important to search for the ease adsorbent. One potential material that can be created as a modest and, effectively accessible is strong waste from the welding business utilizes calcium carbide called calcium carbide buildup. As per Jiang [11], calcium carbide buildup has a high range of pH somewhere in the range of 12.84 and 13.2, explicit gravity of 2.32, explicit surface territory is 24.664 m<sup>2</sup>/g. This examination planned to research the capability of calcium carbide buildup as an adsorbent for diminishing manganese (Mn) content from AMW with the goal that it is appropriate to be discarded to the accepting water body.

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### 2. MATERIALS AND METHOD

#### 2.1. Preparation of the Adsorbent Material

Calcium carbide residue (CCR) from a neighborhood welding in Palembang, South Sumatra, Indonesia was utilized as an adsorbent. In the wake of cleaning and absorbing distillate water medium-term and dried normally in sunshine, and sieved into 80 work of molecule estimate. The adsorbent was made into tablets structure with a

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thickness of 2 mm and width of 3 mm and warmed utilizing the broiler at 150 °C for 120 minutes. The normal load of one tablet dry adsorbent is 0.5 g [12].

Manufactured corrosive mine water was set up by dissolving  $M_n[SO_4]_{4,4}H_2O$  into refined water to acquire a centralization of 100 mg/L and put in a bottle with limit of 40 L at room temperature. To get the ideal focus dilution, a solution of  $H_2[SO_4]_0$ , 1 N was included request to alter the pH of arrangement and filtered using Whatman channel paper No. 1.

## 2.2. Batch Adsorption Experiments

The process of adsorption of Mn(II) ions was carried out using a batch method at room temperature with initial concentrations of 20, 40, 60, 80, 100 mg/L and the weight of adsorbent calcium carbide residue were 2, 5 and 7.5 g, mixed into 100 mL of synthetic AMW solution. The mixture was stirred using shaker with a speed of 200 rpm for 180 minutes and the sample is taken periodically from 5 to 180 minutes.

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## 2.3. Adsorption Capacity and Removal Efficiency

The amount of reduction in Mn(II) ions per mass of adsorbent at a certain time ( $q_t$ ) is calculated based on the formula in Eq. (1) [13].

$$q_t = \frac{(C_0 - C_t) * V}{m} \quad (1)$$

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The percentage of adsorption of heavy metals iron (II) ions from solution is calculated using the following equation [14]:

$$\text{Adsorption (\%)} = \frac{(C_0 - C_t)}{C_0} \times 100\% \quad (2)$$

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Where,  $q_t$  is the adsorbent retention limit at time  $t$ ;  $C_0$  is the underlying grouping of the metal in arrangement (mg/l);  $C_t$  is the convergence of metal in arrangement after time  $t$  (mg/l);  $V$  is the volume of arrangement (L) and  $m$  is the mass of the adsorbent in arrangement (g).

The Langmuir adsorption condition is a standout amongst the most widely recognized isotherm conditions used to show balance information in a strong fluid framework. This condition applies to the monolayer surface with various indistinguishable locales disseminated homogeneously on the outside of the adsorbent. The general type of the Langmuir condition is communicated in a straight structure, as pursues [15, 16].

$$\frac{C_e}{q_e} = \frac{1}{bq_m} + \frac{C_e}{q_m} \quad (3)$$

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Where  $b$  and  $q_m$  are Langmuir constants which are identified with the adsorption rate (L/mg) and adsorption limit (mg/g).

The Freundlich isotherm adsorption model depicts a heterogeneous adsorption process superficially with

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a non-uniform warmth adsorption dissemination [17]. As indicated by the Freundlich isotherm adsorption model, the mass of adsorbate ingested per mass of adsorbent can be communicated as a component of solute fixation,  $C_e$ . The direct type of the Freundlich isotherm adsorption technique is detailed as condition (4):

$$5 \quad \log q_e = \log K_f + \frac{1}{n}(\log C_e) \quad (4)$$

Where  $K_f$  and  $n$  are Freundlich constants. To decide  $K_f$  and  $n$ , every one of them utilizes a direct relapse estimation of the capture and incline diagram ( $\log q_e$ ) versus ( $\log C_e$ ) while the connection R2 gives a sign of the model picked [18].

By utilizing the straight condition come nearer from the chart of Langmuir and Freundlich isotherms, we can know the adsorption limit of CCR adsorbent. Assurance of Langmuir adsorption isotherm was gotten by making a bend of the connection between the harmony focus in the fluid stage ( $C_e$ ) to the strong stage balance fixation ( $C_e/q_e$ ), for Freundlich adsorption isotherm acquired by charting the connection between  $\log C_e$  with  $\log q_e$ .

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2.4. Adsorption Kinetics

Study of adsorption energy is significant in light of the fact that it can give data about procedure elements, for example, adsorption rate, living arrangement time, and motor parameters. In this way, dynamic investigations helping in assess the appropriateness of materials utilized as potential adsorbents in expelling poisons from arrangements [2]. Exploratory information testing [20] or the adsorption response model should be possible with pseudo first request models and pseudo second request models. The pseudo first request model is the condition model that is most generally utilized for the procedure of adsorption of solids from arrangements. This model is an adsorption speed model dependent on adsorption limit [19]. Ho and McKay [20] clarified that some first-request pseudo 26 editions were utilized to recognize dynamic conditions dependent on the adsorption limit of the crystallization of adsorbate in arrangement. The condition of the pseudo first request model can be displayed as pursues.

$$\log(q_e - q_t) = \log q_e - \frac{k_1}{2,303}t \quad (5)$$

By plotting  $\log(q_e - q_t)$  on  $t$ , the correlation value of  $R^2$  will be obtained and the first order ( $k_1$ ) adsorption reaction rate value from the graph slope obtained. Whereas the 12 pseudo-second-order kinetic model given by [20] is defined as.

$$19 \quad \frac{t}{q_t} = \frac{1}{k_2(q_e)^2} + \frac{1}{q_e}t \quad (6)$$

Where  $q_t$  is the number of metal ions absorbed in equilibrium conditions (mg/g); ( $k_2$ ) is the pseudo-second-order rate constant applied (mg/g/h), plot  $t/q_t$  with  $t$  will give a linear line.

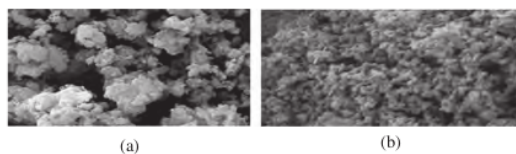
### 3. RESULTS AND DISCUSSION

#### Portrayal of CCR adsorbent

Morphological portrayal of the adsorbent was completed to decide the compound substance and physical and concoction properties of CCR. Portrayal was done utilizing SEM-EDX (Filtering Electron Magnifying lens Dispersive X-Beam Analisis). SEM-EDX is one sort of electron magnifying lens that can create high goals from the picture of an example surface. The portrayal of adsorbent was likewise done by infrared spectroscopy or normally called Fourier Change Infrared (FTIR) which means to decide the utilitarian gatherings of mixes contained in CCR as appeared in Figure 1.

Figures 1(a) and (b), demonstrate that there are morphological changes on the outside of CCR when adsorption process. This demonstrating there is an accelerated of Mn(II) particles in the adsorbent surface. This outcome is likewise upheld by the EDX (Vitality Dispersive X-Beam) investigation which is available the components in the CCR before and after the adsorption process. Characterization utilizing EDX to acquire data on the substance of component<sup>25</sup> found in the CCR, as appeared in the range of portray<sup>30</sup> in Figures 2(a) and (b), and Table I.

From Figures 2(a) and (b), and Table I, it very well may be seen that the CCR before being utilized as adsorbent was dominate by calcium, and after the adsorption procedure the calcium substance diminishes from 52, 83 to



**Fig. 1.** (a) Surface morphology of adsorbent the CCR before adsorption (magnificent of 5000 times). (b) Surface morphology of adsorbent the CCR after adsorption (magnificent of 5000 times).

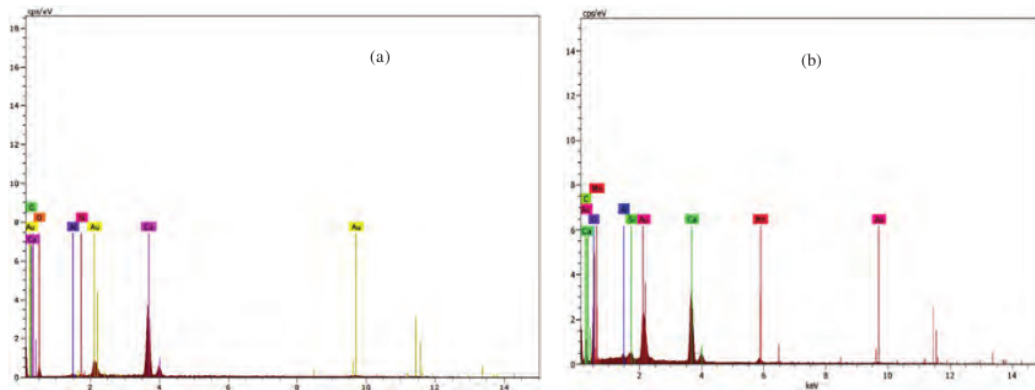
**Table I.** The composition of the CCR before and after adsorption.

Component/composition (%)			
Before adsorption		After adsorption	
Elements	(wt.%)	Elements	(wt.%)
Ca	52.83	Ca	30.95
Au	24.30	Au	25.26
O	16.56	O	28.15
C	3.76	C	8.27
Al	1.56	Al	0.79
Si	0.98	Si	1.01
		Mn	5.57
Total	100	Total	100

30.95 wt%. While the substance of Mn(II) particles exist after the adsorption procedure with a measure of 5.57 wt%.

Portrayal of utilitarian gatherings of the CCR was additionally done by FTIR Spectrophotometer to recognize the natural mixes, both subjectively and quantitatively (Figs. 3(a and b)). The investigation is finished by taking a gander at the state of the range and tops that show the kind of practical gathering controlled by the compound. The consequences of FTIR portrayal of practical gatherings of the CCRs acquired with a recurrence scope of 4000–500 cm<sup>-1</sup> are demonstrated by a diagram of the<sup>10</sup> connection between the wavelength and Transmittance, as appeared in Figures 3(a) and (b).

In view<sup>10</sup> of the consequences of FTIR when adsorption as appeared in Figures 3(a) and (b). A portion of the pinnacles saw in the range demonstrate that the CCRs have a useful<sup>3</sup> gathering crest which can tie cations. At the wavelength of 3637.9 cm<sup>-1</sup> and 3362.1 cm<sup>-1</sup> compares to the extending of the hydroxyl (-Goodness) work within the vibration which shows the nearness of COO-symmetric carboxyl gatherings, suggesting the nearness of kaolinite elements [21, 22]. While at the stature of the band tops 2877.5, 2817.97 and 2512.2 cm<sup>-1</sup> speaking to flexural (-Goodness) vibrations and furthermore as



**Fig. 2.** (a) Characteristic spectrum of CCR before adsorption. (b) Characteristic spectrum of CCR after adsorption.

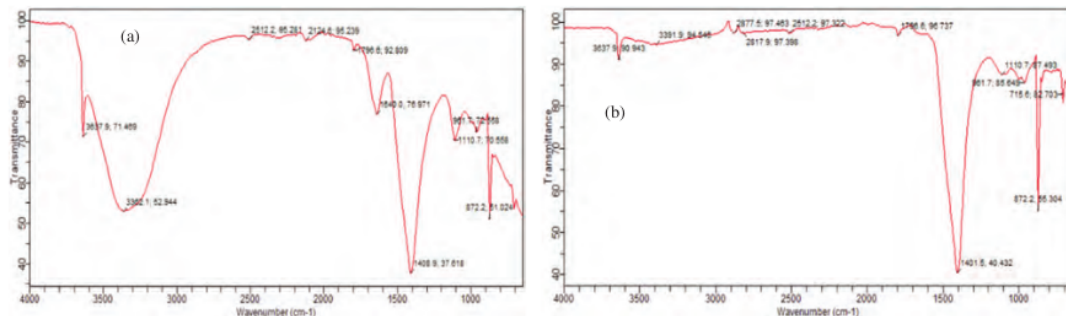


Fig. 3. (a) FTIR of CCR before adsorption. (b) FT-IR of CCR after adsorption.

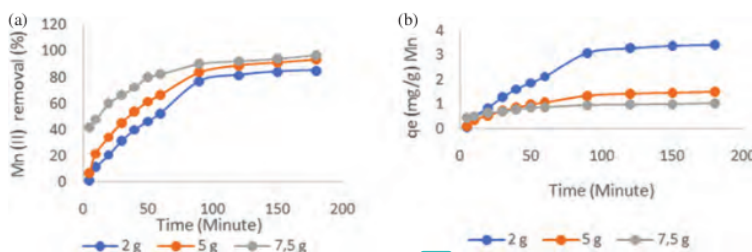


Fig. 4. (a) Effect of mass of adsorbent to percent reduction in Mn(II) ions. (b) Effect of mass of adsorbent to the adsorption capacity of Mn(II) ions.

symmetrical COO extend vibrations, showing the nearness of hydroxyl [23, 24], then sharp crests at 1401.5 and 1110.7  $\text{cm}^{-1}$  were because of Si-O strain vibrations, for retention band statures 961.7 and 83.2 identified with flexural vibrations demonstrating the presence of Al-O gathering [25], and there is another band as high as 715  $\text{cm}^{-1}$  related with vibration Mn-O extending in manganite [26].

### 3.1. Effect of the Adsorbent Period

The information got plot the connection between the measure of metal Mn(II) particles adsorbed to time and plot the measure of metal expulsion of Mn(II) particles to the season of every time of the CCR, as appeared in Figure 4.

Figure 4(a) the level of the CCR take-up is exceptionally quick in expelling Mn(II) particles and increments with expanding mass of the CCR. The adsorption limit of the CCRs achieves immersion stage at an hour and a half nearly for all mass of carbide calcium buildup. The underlying reduction of Mn(II) is quick because of the expansion in the outside of the adsorbent and along these lines delivers a progressively dynamic site for Mn adsorption. Figure 4(b) demonstrates the impact of contact time and adsorption limit of the CCR. During the principal hour, the adsorption limit demonstrated a fast increment and afterward gradually expanded until it achieved the ideal time at 100 minutes. The level of evacuation is higher toward the beginning of this trial because of the accessible

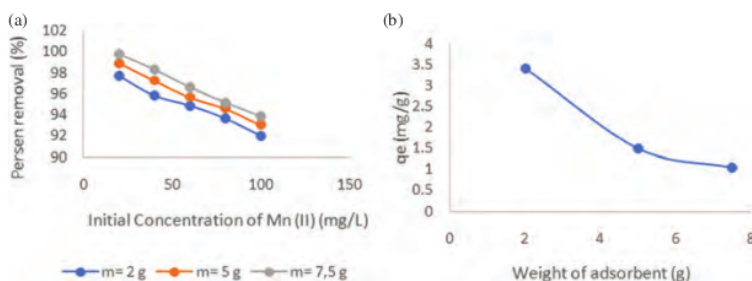


Fig. 5. (a) Effect of initial concentration on percent adsorption of Mn(II). (b) Effect of mass of adsorbent on absorption power.

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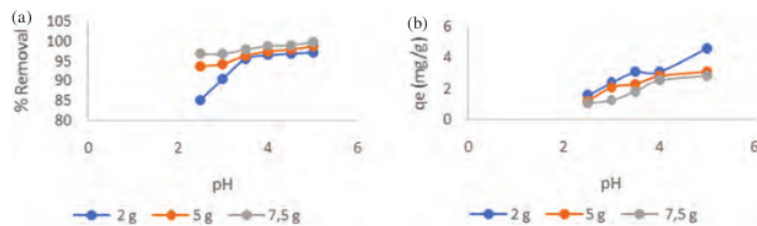


Fig. 6. (a) Effect of pH on the adsorption percentage. (b) Effect of absorption rate on pH.

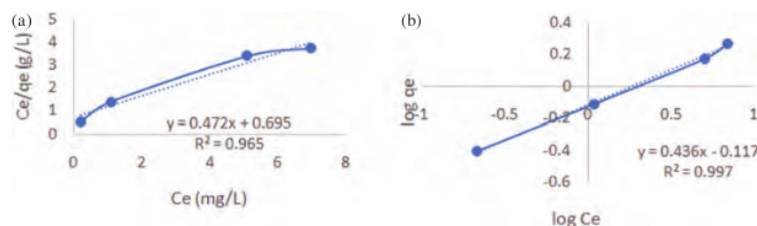


Fig. 7. Langmuir and Freundlich plot graphs for Mn adsorption with adsorbent calcium carbide residues. (a) Langmuir model. (b) Freundlich model.

surface territory of the CCR for retention of metal Mn(II) particles.

### 3.2. Effect of Mn(II) Ion Concentration

The impact of beginning convergence of Mn(II) arrangement was examined with different mass carbide calcium buildups, in particular 2, 5 and 7.5 gr. The calcium carbide buildup put into 100 ml of Mn(II) arrangement with fluctuating fixations, between 20 to 100 mg/L with an unsettling rate of 200 rpm. The impact of the underlying fixation on the adsorption of Mn(II) particles utilizing the CCR for 180 minutes is appeared in Figure 5.

In Figures 5(a) or (b), it tends to be seen that the level of adsorption of Mn(II) particles diminishes altogether with an expanding beginning convergence of Mn(II) particles in arrangement. At the underlying centralization of 20 mg/L for the mass of adsorbent carbide calcium buildup 2 gr, the level of adsorption of Mn(II) was 97.7% diminishing to 91.98% at the underlying convergence of Mn(II) particles of 100 mg/L. While for the mass of adsorbent carbide calcium buildup 5 gr, there was a reduction from 98.93% to 93.03%. For mass, the leftover carbide calcium is 7.5 gr, the level of adsorption of Mn(II) particles drops from 99.8% to 93.9%. In Figure 7(b) it tends to be seen that the more noteworthy the mass of the adsorbent,

the more prominent the capacity of lingering calcium carbide to adsorb metal Mn(II) particles.

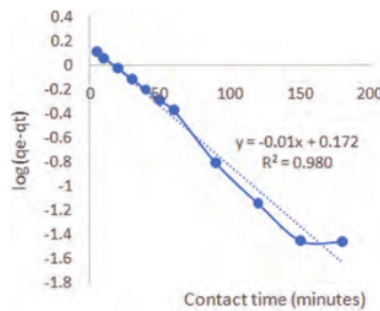
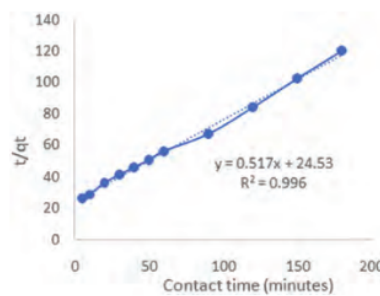
### 3.3. Effect of pH on Mn(II) Ion Removal

The effect of pH in this study was studied in the range of 2.5; 3.5 and 5 at room temperature. The initial concentration of Mn(II) ion solution was set at 80 mg/L, stirring speed 200 rpm, contact time of 160 minutes and mass variation of adsorbent 2, 5 and 7.5 g. The results of the analysis obtained are shown in Figure 6.

In Figure 6 it very well may be seen that the expansion in the underlying pH of the arrangement can build the measure of Mn(II) particles consumed by the remaining carbide. The level of manganese expulsion at pH 2.5–5 is moderately high at 97.25% for the adsorbent load of 2 g, 98.73% for mass adsorbent 5 g and 99.9% for mass adsorbent 7.5 g. While the adsorption limit ( $q_e$ ) of Mn metal by the most extreme carbide calcium buildup happens in the mass of 2 gr adsorbent with an underlying pH of 5.0 arrangement, which is equivalent to 5.6 mg/g and the least at 7.5 g adsorbent mass with the underlying pH of arrangement 2, 5, which is equivalent to 1.03 mg/g. In these conditions, the pH of the blend in the adsorption procedure goes up to 10, 5. This is because of the exceptionally high pH of the underlying carbide

Table II. Parameters of adsorption of Mn ions as a result of linearization of the equation of the Langmuir and Freundlich adsorption models.

Time (minute)	Langmuir					Freundlich				
	Linear equations	$q_0$	$R^2$	$K_L$	$R_L$	Linear equations	$R^2$	$K_F$	$1/n$	
180	$Y = 0.4723X + 0.695$	2,117	0,9652	0,223	0,01	$Y = 0.4368X - 0,1173$	0,997	0,763	0,437	

Fig. 8. Relationship  $\log(q_e - q_t)$  curve versus  $t$ .Fig. 9. Relationship curve  $t/q_t$  versus  $t$ .

calcium buildup, which is 12.5. At the high pH estimation of the blend, the grouping of hydrogen particles as contenders is lower, causing an expansion in the quantity of metal particles Mn(II) assimilated from the arrangement [27, 28]. At low pH beneath 4 the quantity of H<sup>+</sup> particles are extremely enormous so that there is rivalry between Mn(II) particles and Hydrogen particles to keep the dynamic adsorbent gathering, this challenge causes disturbance of the retention procedure of Mn(II) particles with the goal that the measure of substantial metal particles adsorbed is low [28].

### 3.4. Equilibrium Adsorption Isotherms

Model Langmuir and Freundlich adsorption isotherm are utilized to depict a gathering of information acquired at a specific fixation scope of perceptions. The perceptions in this examination led by the underlying centralization of the particle Mn(II) 80 mg/L and mass of adsorbent remaining calcium carbide 5 g contact time 180 minutes.

To decide the suitable isotherm condition on research by taking a gander at the relationship coefficient ( $R^2$ ) got from the direct diagram of every condition. A relationship coefficient near one can be said to pursue the sort of adsorption isotherms of the adsorption isotherm condition, as appeared in Figure 7.

From Figure 7 the price of the Langmuir and Freundlich constants can be calculated from the linear  $C_e/q_e$  versus  $C_e$  plot and  $\log q_e$  versus  $\log C_e$ . As shown in Table II.

The most extreme adsorption limit ( $q_o$ ) of Table II above, at the contact time of 180 minutes as per the Langmuir model is 2.117 mg/g and the  $R^2$  worth is 0.9652 and the consistent cost of Langmuir partiality ( $K_L$ ) is 0.223 L/mg and ( $R_L$ ) of 0.01 (generally excellent) on the grounds that it is littler than 1 ( $0 < R_L < 1$ ). While for the Freundlich isotherm model the cost of  $R^2$  is acquired at 0.9974 and the cost of Freundlich consistent ( $K_F$ ) is 0.763 and the force esteem ( $1/k_F$ ) is 0.4368. From Table II, it very well may be presumed that the adsorption isotherm model in Figure 7 is the grouping of Mn(II) particles pursues the Freundlich isotherm model. The Freundlich adsorption isotherm model demonstrates a direct connection between the quantity of substances adsorbed per gram of adsorbent in the adsorption procedure of Mn(II) by the CCR, with the condition  $q_e = 0.736C_e^{0.4368}$ , this implies the isotherm design in engrossing Mn metal particles (II) by the adsorbent of calcium carbide strong waste, because of the dynamic side of the strong waste surface calcium carbide is multilayer, with irregular dispersion and liking over heterogeneous surfaces.

### 3.5. Study of Adsorption Kinetics

The dynamic model utilized in this examination is pseudo-first request, pseudo-second request. Both of these dynamic models are ordinarily used to decide the request of adsorption energy [19]. To discover how the system, viability and productivity and how quick the adsorption procedure of Mn(II) particles happens utilizing adsorbent the CCRs, it is important to think about pseudo-first-request and two-pseudo request energy. Perceptions were done at the underlying convergence of 80 mg/L and the mass of adsorbent carbide buildups 5 g with contact times of 5 to 180 minutes, information acquired by communicating conditions 4 and 5. Insertion after effects of adsorption of metal particle Mn(II) between  $\log(q_e - q_t)$  to  $t$  appeared in Figure 8 and between  $t/q_t$  versus  $t$  in Figure 9. The parameter esteems and connection coefficients got

Table III. Parameters and linearity equation of one and two pseudo-order kinetics models.

Massa RCC	Pseudo kinetic model first order				Pseudo kinetic model second order			
	Linear equation	$q_e$	$R^2$	$k_1$	Linear equation	$q_e$	$R^2$	$k_2$
2	$y = -0.0158x + 0.8824$	7.628	0.911	0.125	$y = 0.1869x + 17.653$	5.350	0.9920	0.0019
5	$y = -0.01x + 0.1724$	1.487	0.981	0.023	$y = 0.5179x + 24.532$	1.931	0.9962	0.0109
7,5	$y = -0.0119x - 0.0726$	0.846	0.943	0.027	$y = 0.8622x + 18.326$	1.159	0.996	0.0406

from counts dependent on Figures 8 and 9, as in Table III.

From the relationship coefficient esteems are displayed in Table III. In light of the coefficient of assurance ( $R^2$ ) got from the third time of the adsorbent remaining calcium carbide is connected, the mass of 5 grams of the most great. The estimation of the relationship coefficient ( $R^2$ ) most near 1, i.e., = 0.9962 by the quantity of particle Mn(II) adsorbed at harmony ( $q_e$ ) is 1.931 mg/g, with the pace of the energy of  $k_2$  to be specific 0.0109, gr. mg/minute.  $R^2$  qualities got from particle adsorption of Mn(II) by the CCR following the model of the pseudo-second-request energy. As indicated by Defeat [29], and Huang [30] when adsorption models pursue the model of pseudo-second-request energy, it tends to be sorted adsorption process which happens between the CCR and particle Mn(II) is synthetic adsorption (chemisorption) [31–33].

#### 4. CONCLUSION

Adsorption limit and adsorption proficiency are expanded with expanding pH<sup>6</sup> and diminished with expanding in weight of adsorbent. The adsorption limit of the outside of the CCRs for Mn(II) particles increments with an expansion in pH and introductory convergence of the arrangement of Mn(II) ions [34].

Testing the Freundlich and Langmuir isotherm models for test information demonstrates that: the Freundlich isotherm model matches the exploratory outcomes. The Freundlich adsorption isotherm model demonstrates a direct connection between the measure of adsorbed substance per gr adsorbent in the adsorption procedure of Mn(II) with the condition  $q_e = 0.736C_e^{0.436}$ .

The use of pseudo-first-request energy and <sup>6</sup>pseudo-second-request dynamic models demonstrates that the adsorption of Mn(II) particles on the CCRs, most intently pursues the second-request pseudo energy model.

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