ABSTRACT
This study was aimed to determine the interaction of cement kiln dust - aqueous solution contaminated with Cadmium ions was studied through set of batch tests with operational conditions of contact time ≤120 min, sorbent dosage from 0.05 to 1 g/100 mL and agitation speed ranged from 50 to 250 rpm for initial concentration of 50 mg/L with initial pH of 3 to simulate the acetogenic phase in the sanitary landfill. The best values of these conditions are 1 hr, 0.7 g/100 mL and 250 rpm respectively, To obtain maximum removal efficiencies of 97.6%. Freundlich and Langmuir models have a high ability in the representation of the sorption data with determination coefficient (R^2) greater than 0.97 and the sorption capacity reached to 84.1 mg/g. This certifies that the physical sorption and chemisorption can occur together to remove Cadmium ions from the aqueous solutions. Tests elucidated that the average coefficient of the hydraulic conductivity is equal to 9.7×10^{-13} m/s and this is suitable for LPB. Finally, COMSOL Multiphysics 3.5a package was able to simulate the distribution of cadmium ions concentrations within two-dimensional physical model packed with sand aquifer and CKD LPB. A good matching between model predictions and experimental results are recognized at selected points up and down gradient of LPB.

Keywords: barriers technology; groundwater contamination; COMSAL software; sorption.
INTRODUCTION
Water present beneath the ground surface is known as “groundwater” which can be infiltrated from the surface by migration through the soil matrix of geologic formations. It becomes necessary to exploit the groundwater. Arab World has a groundwater reserve around 14.3 thousand billion cubic meters (11). Groundwater is a vital source because previous records signified that the 2 billion people depends mainly on this water for domestic uses and the 40% of food production through the worldwide is based on irrigation with groundwater; accordingly, this water considers a vital resource (18,21). Previous studies signified that the contamination of aquifer will prohibit the use of its water for long periods of time reached to several decades (17). Heavy metals like Cadmium have a good ability for accumulation in the web of aquatic food to reach the human beings and cause many pathologies (2). In Japan, the spreading of Itai-itai disease in the 1950s was attributed to the presence of Cadmium element (resulted from base-metal mining) with high concentrations in water applied in the irrigation of rice fields. The heavy metals in groundwater may be resulted from water infiltration through the polluted layers of the soil (9). Also, solid wastes like municipal solid waste incineration ash, waste rock and others may be represented a familiar sources of the heavy metals through the leaching process. The groundwater must be protected by applying the familiar technologies like permeable reactive barriers (PRBs), low permeable barriers (LPBs). The LPBs technology can be used to contain contaminants in an aquifer and prevent the further spreading of contaminants in groundwater. Applying this technique in the restriction of groundwater contaminated with heavy metals has been proved to be successful and most efficient in the redirection or capturing of contamination plume. The heavy metals can be captured by the reactive material used within the low permeable barrier technology by the following mechanisms: chemical precipitation, sorption, ion exchange, abiotic reduction and biotic reduction. In fact, with many cases, remediation of contaminated groundwater can be achieved by one or more of these processes. To keep LPB costs to the minimum, the material should be persisting with the time and readily available at low cost. Cement kiln dust (CKD) is alkaline powder solid waste resulted from the cement industry as byproduct; however, about 41 kg of this waste can be accompanied with production of 1-ton of cement (23); so, huge amounts of this waste will form a burden on the environment (4). The popular material used in the restriction the contaminant transport in the subsurface environment is the clay (or bentonite) as in the liner of the sanitary landfill to prevent the movement of leachate or in the hydraulic barrier for redirection the flow faraway of water resource. So, the reuse of CKD as an alternative choice to the bentonite due to its mentioned information is very important in the fields of sustainable development and green projects as well as the reducing of disposal charges (15). The CKD is classified chemically as a high heterogeneous material and, therefore, this material can remove the dissolved metal in the aqueous solution by adsorption and precipitation (22). The relationships between water content, compaction, hydraulic conductivity, and dry unit weight have identified in the rigid- and flexible- wall permeameters. The results proved that the suitability of the sand in the formation of low hydraulic conductivity barrier layers to be used as pits and covers in landfills, ponds, and sewage lakes (3). The 2D steady groundwater flow beyond the LPB embedded in the flow domain was solved analytically for finding the discharge vector and hydraulic head. Also, the developed solution applied to investigate the role of extraction wells and vertical barrier in the containment of pollution front (6). The ability of synthetic clay linings (GCL) to prevent contaminants from moving into groundwater as they are used in the design of the landfill barrier was studied (8). The study was implemented to investigate the applicability of seven clayey substrates taken from countryside regions in Poland as durable, successful and sustainable materials for constructing the liners of the waste landfill for preventing the leachate seepage to surrounding environment (16). This study was aimed to consider a good application for the principles of sustainability
because it investigates the possibility of using cement kiln dust solid waste as LPB as alternative to natural clay for preventing a further spreading of Cadmium ions in the groundwater. The performance of this material was evaluated by sets of batch and continuous tests as well as COMSOL Multiphysics 3.5a (2008) used to simulate the spreading of the Cadmium ions.

**MATERIALS AND METHODS**

**Materials:** The CKD was collected from the Al-Kufa cement factory, Kufa, Al-Najaf city, Iraq with physical appearance shows in Fig. 1. It considers heterogeneous material with composition consisted of CaO, SiO₂, Al₂O₃, Fe₂O₃, MgO, SO₃, Na₂O and K₂O with percentages of 43.99, 15.05, 4.43, 2.23, 1.64, 6.02, 0.69 and 4 (14) (25). The specific surface area of CKD was found equal to 5200 cm²/g measured according to ASTM C 204-05 using the Blaine-air permeability apparatus available in the Physical Laboratory of Kufa Cement Plant, Southern Cement Company (20). Sand was used, sieved for size ranged from 0.6 mm to 1 mm. This quartz sand was obtained from the granular filter factory in Najaf, Iraq. The sand was washed thoroughly with distilled water before use and then dried using the oven for 2 hours under 230°C. Quartz sand is characterized by a uniform distribution of grain size with median size (d₅₀) of 0.7674 mm. The bulk density, porosity and hydraulic conductivity coefficient have value of 1.7335 g/cm³, 0.31, and 4.1×10⁻⁴ m/s respectively.

**Contaminant:** To prepare the water sample with Cadmium concentration of 1000 mg/L, 2.744 g of Cd(NO₃)₂·4H₂O was dissolved in one liter of distilled water and this solution was kept at room temperature. The solution was used as stock solution to prepare any specific concentration of Cadmium by dilution and its pH adjusted to became 3 to represent the acetogenic phase of leachate by adding 0.1 M HNO₃ or NaOH as required.

**Batch experiments:** They were carried out to determine the best conditions that required to achieve the maximum removal efficiency of pollutant. These conditions included contact time, adsorbent dosage, initial metal concentration and agitation speed for certain value of initial concentration and initial pH of the solution. Series of 250 mL flasks were employed and each flask was filled with 100 mL of cadmium solution which has initial concentration (C₀) of 50 mg/L. The CKD dosages (m) of (0.05, 0.1, 0.3, 0.5, 0.7, 0.9 and 1 g) were added to the different flasks and they have been stirred in the high-speed orbital shaker (Edmund Buhler SM25, German) at 250 rpm for 1 hour. A fixed volume (20 mL) of the solution was withdrawn from each flask and filtered using filter paper type (JIAO JIE 102) to separate the adsorbent; then, fixed volume (V =10 mL) of the clear solution was utilized for the determination of the amount of unsorted metal ion still present in solution (Cₑ). The measurements were carried out to concentration of cadmium using atomic absorption spectrophotometer (AAS, Sens AA, Australian). Different values of contact time (0, 50, 100, 150, 200 and 250) and agitation speed (50, 100, 150, 200 and 250 rpm) for initial pH of 3 and initial concentration of 50 mg/L. The quantity of adsorbed contaminant (qₑ) was established by the mass balance using the following equation (10):

\[
qₑ = (C₀ - Cₑ) \frac{V}{m}
\]  

Figure 1. Physical appearance of cement kiln dust used in this study
The removal efficiency ($R$) for all contaminants was determined by:

$$ R = \left( \frac{C_o - C_e}{C_o} \right) \times 100 \quad (2) $$

**Pure precipitation**

In the present study, a set of batch experiments for finding the precipitation portion that occurs due to interaction of CKD and Cadmium ions have been conducted. The results of these experiments with outputs of experiments conducted to describe the total sorption can be used together to find the contribution of pure adsorption and pure precipitation in the total sorption of removal process. Batch equilibrium experiments for finding the removal of cadmium ions by pure precipitation were carried out by adding certain mass of CKD sorbent into 100 mL of acidic uncontaminated water ($\text{pH} = 3$) and the solution was kept stirred at high-speed using orbital shaker for 1 hr at 250 rpm. The solution was then filtered to separate the sorbent using a procedure similar to the "tea bag" experiment (7). The clear solution was mixed with aqueous solution contaminated with cadmium ions and the resulting solution was filtered after the required equilibrium time. The concentration of the metal ions still present in the filter solution was measured by AAS and the precipitated concentration was obtained by the mass balance.

**Continuous experiments**

The 2D tank (Fig. 2) was performed to represent the migration of contaminant in the presence of CKD LPB embedded within the packed sandy aquifer. This glass tank has dimensions of 100 cm × 40 cm × 10 cm (L × W × H) and it is divided into three compartments by inserted two vertical perforated glass plates. These plates was covered with filter papers, provided the lateral boundaries for middle compartment (with dimensions 80×40×10 cm) which is the location of the packed bed consisted of sand aquifer and CKD barrier. Storage tank with capacity of 80 L, constant head reservoir, and flow meter are utilized to supply the required flowrate for packed bed. Samples have been taken periodically from the flowing water through the model aquifer by using stainless needles. Firstly, the 5 cm bed depth was packed in the middle compartment in the alignment measured from left side of the tank and composed of 50 cm sandy soil, 10 cm CKD LPB, and then 20 cm of the sandy soil. Monitoring of Cadmium concentrations in the selected sampling ports as shown in Fig. 2 for duration not exceeded 14 days was achieved. The 2 mL water samples were taken regularly (after 2, 4, 8…. 336 hours) from the mentioned ports to measure the cadmium concentration by AAS.

![Figure 2. Set-up of two-dimensional experimental tank used in the present study](image-url)
Modeling of batch and continuous outputs

Models of sorption

They related between the contaminant concentration sorbed on the solid phase of the sorbent \( q_e \) and concentration remaining in the liquid phase \( C_e \). The sorption models have many mathematical relationships like Langmuir and Freundlich as below:

\[
q_e = \frac{q_{\text{max}} b C_e}{1 + b C_e}
\]

where \( b \) and \( q_{\text{max}} \) are the affinity constant (L/mg) and the maximum adsorption capacity (mg/g) respectively.

Freundlich model can be written as follows:

\[
q_e = K_F C_e^{1/n}
\]

where \( K_F \) is the Freundlich constant and \( 1/n \) is the intensity of sorption \( (12) \).

Modeling of Cadmium transport

The model developed in this study to simulate the transport of cadmium ions in the 2D bed is based on the combination of two main equations; the first one for the flow of groundwater (Eq.5) and the second equation describes the transport of contaminant (named advection-dispersion equation, Eq.6) as follows:

\[
\frac{\partial}{\partial x} (K_x \frac{\partial h}{\partial x}) + \frac{\partial}{\partial y} (K_y \frac{\partial h}{\partial y}) = 0
\]

\[
D_x \frac{\partial^2 C}{\partial x^2} + D_y \frac{\partial^2 C}{\partial y^2} - V_x \frac{\partial C}{\partial x} = \frac{\partial C}{\partial t} + \rho_b \frac{\partial q}{\partial t}
\]

where \( K \) is the hydraulic conductivity coefficient (m/s), \( D \) is the dispersion coefficient (m\(^2\)/s), \( V \) is the seepage velocity (m/s), \( h \) is the hydraulic head (m), \( \rho_b \) is the bulk density (kg/m\(^3\)). The Langmuir model must be incorporated with the last equation to describe the adsorption process in the transport of the contaminant and the resultant equation can take the following form:

\[
D_x \frac{\partial^2 C}{\partial x^2} + D_y \frac{\partial^2 C}{\partial y^2} - V_x \frac{\partial C}{\partial x} = \frac{\partial C}{\partial t} + R \frac{\partial q}{\partial t} \tag{7}
\]

and \( R \) will be:

\[
R = 1 + \frac{\rho_d}{n} \left( \frac{q_{\text{mb}}}{1 + b C_e} \right)^2 \tag{8}
\]

where \( R \) is the retardation factor.

RESULTS AND DISCUSSION

X-ray diffraction analysis

The chemical composition of CKD sorbent was characterized using the X-ray diffraction in Germany Laboratory. Geology Department. College of Science University of Baghdad. Fig.3 shows the XRD for CKD and it is clear that the Calcium and Silica oxides are the major constituents of its structure. All calcium in the CKD is assumed to be in the form of calcium oxide (CaO); however, this not the real case. Some of the calcium may be in the form of Calcium carbonate (CaCO\(_3\)) (i.e. unreacted limestone), or Calcium sulphate (CaSO\(_4\)). The presence of lime is measure of the CaO content that readily available for reactions. Also, there is high content of free CaO (free lime), Calcium hydroxide, sulfates, and total alkalis. So, the CKD tends to have higher alkalinity behavior due to the mentioned constituents which play a potential role in the precipitation of the heavy metal. The presence of particles other than Calcium oxide can affect the solubility of CaO into Ca(OH)\(_2\). Specifically, interfering oxides such as magnesium oxide, silic, aluminum oxide, and unreacted CaCO\(_3\) particles may slow the rate of Ca(OH)\(_2\) dissolution. Oxides of potassium and magnesium can react with water to form the hydroxides of these elements which able to neutralize of acids in environmental systems.

Figure 3. The X-ray diffraction (XRD) analysis for CKD
Coefficient of hydraulic conductivity

The “falling head permeameter” utilized in the measurement of the hydraulic conductivity coefficient for bentonite material. The experiment was beginning by saturating the porous medium sample with tap water for duration not less than 2 days. The test is based on the monitoring of the falling head as a function of the time and the corresponding coefficient of hydraulic conductivity can be calculated. Results proved that the coefficient of hydraulic conductivity have the values of was decreased with the time from $2.05 \times 10^{-12}$, $1.30 \times 10^{-12}$, $9.20 \times 10^{-13}$, $5.82 \times 10^{-13}$ and $3.22 \times 10^{-13}$ cm/s after 1, 3, 5, 7 and 9 days respectively with average value of $3.22 \times 10^{-13}$ cm/s. Due to the high density of the present reactive materials, this will cause the slowed flow within the internal pores and, consequently, the coefficient of hydraulic conductivity ($K$) will be low (24).

Batch experiments

Sorbent dosage: The relationships between the metal ions removal efficiencies and sorbent dosages of CKD are plotted in Fig. 4(a) for condition specified within this graph. It can be seen that the increase of sorbent mass will improve the removal efficiency and this may be attributed to available greater vacant sites. Also, this figure certifies that the sorption efficiency was increased from 48 to 90% due to increase of dosage from 0.05 to 0.3 g/100 mL. Beyond the 0.3 g/100 mL, slow increase in the efficiency can be recognized; so, the dosage was chosen of 0.7 g/100 mL to obtain of the fair value of removal efficiencies for Cadmium ions that reached to 97.6%.

Contact time: Fig. 4(b) plots the variations of sorption efficiencies for Cadmium ions onto CKD versus the contact time under the relevant conditions specified within this graph. This figure reveals that the sorption rate is very fast within the first five minutes and at which the efficiency approaches to 94.05%. Thereafter, the rate was gradually slowed and efficiency approximately stabilized beyond 1 hr at 97.6%. The presence of sufficient numbers of the binding sites on the adopted sorbent may be the main reason for high rate of sorption process and its decrease with elapsed time can be resulted from the occupation of these sites by cadmium ions; however, the next tests can be implemented with contact time of 1 hr.

Agitation speed: This speed was changed within the range extended from 50 to 300 rpm and the measured corresponding removal percentages of the cadmium ions have been plotted in the Fig. 4(c) under the operational conditions of $C_0= 50$ mg/L, contact time= 1 hr, initial pH= 3, and dosage= 0.7 g/100 mL. For CKD, the slight increase in the sorption efficiency from 91.8 to 100% can be observed due to change of agitation speed from 50 to 300 rpm. This means that the reactivity of CKD is very high due to generate two mechanisms namely adsorption and precipitation responsible of the removal process; so, the agitation speeds within the adopted range not have significant effect on the removal process. For practical purposes, the choice speed of 250 rpm as best value for batch tests is satisfactory for two sorbents under consideration.
Figure 4. Trend of the relationships related between the removal efficiencies of the cadmium ions a) sorbent dosage, b) contact time and c) agitation speed

Pure precipitation in the Cadmium ions-CKD interaction

The removal of metal ions due to interaction with CKD sorbent is governed mainly by two mechanisms pure adsorption and pure precipitation. Based on the total sorption and pure precipitation measurements with applying the same procedure adopted by (1), the proportions of cadmium ions removed by pure adsorption and pure precipitation are plotted in Fig.5. In comparison with total sorption findings, it seems that there is a lag in the removal of cadmium by pure precipitation because the metal ions for this case not in direct contact with CKD. Also, this figure proved that the increase of sorbent dosage will associate with increase of metal ions removal due to the pure precipitation and, consequently, the decrease of ions removed by pure adsorption. This figure showed that the pure precipitation is governed the removal process in the high Cadmium concentrations due to the formation of Cadmium hydroxide as a result of the chemical reaction of CKD. In addition, the precipitation process may be faster than the adsorption process and this requires relatively longer time to achieve the maximum adsorption capacity. Accordingly, precipitation represents the predominant mechanism at high concentrations of the removal process, while adsorption will be the predominant mechanism at low metal concentrations.
Figure 5. Pure adsorption and pure precipitation shares for removal of Cadmium ions onto CKD sorbent

Isotherm models for sorption data

Fig. 6 shows the isotherm models (Freundlich and Langmuir) that fit the sorption data to describe the removal of cadmium ions onto CKD reactive material. Constant of these models are listed in Table 1 and they are calculated by using non-linear regression fitting in Excel (2016) utilizing the “Solver” option. Results certified that the two models are able to represent of sorption data for case under consideration. The maximum adsorption capacity of CKD is equal to 84.1 mg/g as calculated from Langmuir model with affinity constant of 0.04360 L/mg.

![Isotherms models for sorption of cadmium ions onto CKD](image)

Figure 6. Isotherms models for sorption of cadmium ions onto CKD

| Model | Parameter | Value |
|-------|-----------|-------|
| Freundlich | $K_f (mg/g)(L/mg)^1/n$ | 5.27608 |
| | $n$ | 1.51359 |
| | $R^2$ | 0.98807 |
| Langmuir | $q_{max}$ (mg/g) | 84.1 |
| | $b$ (L/mg) | 0.04360 |
| | $R^2$ | 0.97471 |

Fourier transform infrared (FTIR) analysis: The pattern of Cadmium ions sorption onto the CKD is attributable to the active groups and bonds present on this sorbent material (13). The FTIR spectroscopy may be, therefore, done for specifying the preliminary functional groups present in the native CKD used as sorbent for cadmium ions.
The characteristics bands of the CKD before and after Cadmium uptake at pH 3 were used to assess the changes in the functional groups. Functional groups analysis using the FTIR was measured for samples obtained from the experiment before and after cadmium adsorption where results have been plotted in Fig.7. Within the range of wavelengths from 400 to 4000 cm\(^{-1}\), the Si-O functional groups were observed at the 1107 cm\(^{-1}\). Bands at 3450 and 1450.0 cm\(^{-1}\) indicated the possibility of hydration by water or H-O-H bending of water in the adsorbent (5). FTIR spectra of cadmium ions showed that the peaks at 3450, 1450, and 1140 cm\(^{-1}\) had been shifted due to interaction with Cadmium ions. These shifts may be attributed to the changes in counter ions associated with carboxylate and hydroxyxlate anions which contributors in metal ion uptake (19).

![FTIR analysis for CKD before and after sorption of cadmium ions](image)

**Figure 7.** FTIR analysis for CKD before and after sorption of cadmium ions

Contaminant transport in the 2D bed consisted of aquifer and LPB

Application of COMSOL Multiphysics was achieved to simulate the 2D steady flow of water and transient contaminant transport through the bed identical to the experimental setup mentioned previously. The model has dimensions of 0.8 m length, and 0.4 m width for bed height of 5 cm; also, the alignment of LPB (0.1 × 0.3 m) and line source of 0.10 m. Tables 2 and 3 listed all information required for modeling process. The ability of COMSOL Multiphysics 3.5a in the description of the Cadmium ions transport within the sand aquifer containing the CKD LPB was evaluated by comparison the predictions of the theoretical model with experimental measurements at selected within the LPB as depicted in Figure 8 for the locations (P1, P4, P5). It is clear that there is a satisfactory agreement between the model outputs and experimental results; also, this figure certified that the proposed barrier play a significant in the protection of locations nearby it in the down-gradient and the contaminant concentration was increased with the time. Due to the presence of matching between model results and measurements, the present simulation based on the COMSOL package can be utilized to predict the spatially and temporally normalized concentrations of Cadmium ions within the physical bed. Accordingly, Fig.9 is introduced the predictions of breakthrough curves for locations (P2, P3, P6, P7) up- and down-gradient of LPB. Figs. 8 and 9 elucidate that the concentrations of contaminant have the lowest values in the locations adjacent to the LPB from down gradient side. Also, the hydraulic head and velocity values were plotted in the flow field under the effect of LPB as shown in Fig. 10. The barrier has ability to create two stagnant points in the up- and down-gradient sides of this wall with significant change in the advection front in the locations nearby the LPB. This figure elucidates that the velocity of the flow is decreased significantly inside the barrier, while a smaller region of reduced speed can be observed outside the LPB. Fig.11 plots the contours of Cadmium normalized concentration across the bed model as predicted by the COMSOL software for various times. The LPB situated beyond the line source of contamination is caused a significant reduction in the rate of advective transport with lateral deviation of the contaminant plume as the streamlines spread to pass the barrier.
Table 2. Constants utilized for modeling of 2D transport of contaminant in the LPB and aquifer

| Bed          | Variable | Value  |
|--------------|----------|--------|
| Sand aquifer | $n$      | 0.31   |
|              | $K$      | $4.1 \times 10^{-4}$ |
|              | $\rho_b$ | 1733.5 |
|              | $n$      | 0.62   |
|              | $K$      | $9.7 \times 10^{-15}$ |
| LPB          | $\rho_b$ | 1170   |

$n=$ Porosity; $K=$ Hydraulic conductivity (m/s); $\rho_b=$ Bulk density (kg/m$^3$)

Table 3. Conditions adopted in the modeling of Cadmium ions transport for the 2D laboratory scale tank

| Item                                | Location          | Type               | Value      |
|-------------------------------------|-------------------|--------------------|------------|
| Boundary conditions for fluid flow  | Line source       | Hydraulic head (cm)| 5          |
|                                      | Outlet side       | Hydraulic head (cm)| zero       |
|                                      | All other sides   | Zero flux/symmetry | -----      |
|                                      | Line source       | Concentration (mg/L)| 50        |
| Boundary conditions for solute transport | Outlet side | Advective flux     | -----      |
|                                      | All other sides   | No flux/symmetry   | -----      |
| Initial condition for solute transport                  | Concentration (x,y) at t=0 | zero |  |

Figure 8. Predictions of COMSOL in comparison with measurements for Cadmium normalized concentrations in the selected points for sand aquifer with presence of CKD as LPB

Figure 9. Predicted values of cadmium normalized concentrations using COMSOL package in the selected points for sand aquifer with presence of CKD as LPB
The cement kiln dust resulted from cement industry as byproduct proved have a high ability in the restriction of the Cadmium ions transport present in the aqueous solutions. The best values of the operational conditions for material under consideration are equal to 1 hr, 0.7 g/100 mL, and 250 rpm respectively for initial concentration of 50 mg/L to achieve the maximum removal efficiencies of 97.6%. Results revealed that the average hydraulic conductivity of CKD was equal to $9.7 \times 10^{-15}$ m/s and this is satisfied the requirements of LPB.

1- For interaction of Cadmium ions with CKD, Freundlich and Langmuir models are found able to describe the sorption measurements where the values of $R^2$ not less than 0.97 with maximum sorption capacity of 84.1 mg/g; so, the removal process may be governed by the physical and chemical forces.

2- Mathematical model describes in a good manner the flow of water and distribution of the metal ions within the 2D bed to specify the efficacy of LPB in the protection of the water resources. The outputs of this model as calculated by COMSOL Multiphysics 3.5a are observed in good matching with experimental measurements.

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