Investigating the Sorption Behavior of Cesium and Cobalt in Soil Samples

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Two soil samples were collected from El-Dabaa site and denoted as DNS/C (Dabaa natural soil / Calcite) and DNS/Q (Dabaa natural soil / Quartz). They were characterized using X-Ray Fluorescence (XRF) analysis, Fourier transform infrared spectroscopy (FT-IR) analysis and X-Ray diffractometer (XRD) analysis. The soil samples were applied as solid phases for retention of 134Cs and 60Co radionuclides from their aqueous solutions. The effect of different parameters as pH value, contact time, and competing ion on the sorption behavior of 134Cs and 60Co was studied in static conditions. The maximum sorption capacity of DNS/C was 1.16 and 4.28 mg.g⁻¹ for Cs(I) and Co(II) ions, while it was 3.19 and 4.76 mg.g⁻¹ for sorption of both metal ions onto DNS/Q respectively. Different kinetic models were applied to assess the sorption of Cs(I) and Co(II) ions from their aqueous solutions by soil samples. The sorption process followed pseudo-second-order kinetics clarifying that chemisorption could be regarded as the controlling step. The role of temperature on sorption of Cs(I) and Co(II) was studied and the values of thermodynamic parameters (ΔH°, ΔG° and ΔS°) were determined. 134Cs radionuclides exhibited ΔH° values amounting to -14.5416 and -13.0962 kJ.mol⁻¹ for sorption onto DNS/C and DNS/Q samples. These negative values confirmed the exothermic nature of sorption process. The enthalpy change attained the values 25.3895 and 24.67368 kJ.mol⁻¹ for 60Co radionuclide sorption onto the soil samples DNS/C and DNS/Q. These positive values confirmed the endothermic nature of the sorption process.

Keywords: Radionuclides, Sorption, Soil Samples

Introduction

Pollution of the environment is one of the most important problems facing the world today. This problem is becoming more and more serious and needs to be resolved as quickly as possible due to its importance for the quality of the environment and the health of people. Radioactive pollution is an important problem attracting the attention of many scientists around the world. This is due to the growing demand for nuclear technology and the health risks accompanying with its widespread application. Egypt has decided, at present, to produce electricity utilizing nuclear power reactors. El Dabaa site was selected for the construction of Egypt’s first nuclear power plant; hence, there is an extreme need to obtain information on the sorption capabilities of soil samples in this area towards different radionuclides.

Radioactive waste contains a large amount of radionuclides that are dangerous for human life and environment [1, 2]. Radionuclides of cesium and cobalt are constituents of the radioactive waste streams as fission and activation products, respectively. [3,4]. 134Cs (T1/2 = 2.06 y) that is formed as one of the common fission products of uranium-235 and other fissionable isotopes in nuclear power reactors. It constitutes...
a large problem because cesium has a high water solubility and easily spreads into the surrounding environment. Cesium mobility can be considerably reduced by sorption of its ions on soil and surrounding minerals [5]. $^{60}$Co is a short-lived radioactive isotope produced in nuclear reactors. It can't be produced by nuclear fission. It was produced in small amounts as a result of interaction of neutron with structural materials of the reactor as well as during the routine operation of nuclear power plants. Small quantities can be also released as contaminants in the cooling water or as radioactive waste. It is a corrosive and toxic element widely observed in nuclear effluents and seriously affects the environment [6]. For radiological safety, it is important to remove the radioactive constituents from waste streams and preserve them a stable from. There are many methods that have been successfully applied in removing different radionuclides from hazardous wastes. These methods include flotation [7], common precipitation [8], liquid-liquid extraction [9], membrane processes [10], electrocoagulation [11], adsorption [12] and solid phase extraction technology [13, 14]. The aim of the present work is to explore the sorption capabilities of some soil samples collected from north coast of Egypt at El-Dabaa site towards Cs(I) and Co(II) ions under different experimental conditions.

**EXPERIMENTAL**

**Site of the study**

Nine soil samples and three seawater samples were collected from El-Dabaa site in Egypt that is located on the Mediterranean Sea coast of the North Western desert and lies between longitudes (28°22’ - 28°31’) E and latitudes (31°02’ - 31°05’) N that is about 400 km from the capital (Cairo) and 150 km west of Alexandria. The site is 15 km long and 4 km wide as illustrated in Figure (1). Only, two soil samples were selected and included tin this study.

**Chemicals and reagents**

All chemicals and reagents used in this work were evaluated analytically and used without further purifications. Hydrochloric acid (HCl), sodium hydroxide (NaOH), sodium chloride (NaCl), calcium chloride (CaCl$_2$·6H$_2$O), magnesium nitrate (Mg(NO$_3$)$_2$·6H$_2$O), manganese chloride (MnCl$_2$·4H$_2$O) and aluminum chloride (AlCl$_3$·6H$_2$O ) were purchased from Sigma–Aldrich (USA).

![Fig. (1): Location of study area, El Dabaa, Northwestern Coast, Egypt](image)

**Preparation of radiotracers**

$^{134}$Cs and $^{60}$Co radioisotopes were prepared by neutron irradiation of 0.01 g of cesium chloride and cobalt chloride. The irradiation process was conducted in the Egyptian research reactor in Inshas. Each target was wrapped in a thin plate, placed in a radiation tray, and exposed to a neutron flux of $10^{14}$ nm$^{-2}$.s$^{-1}$. The irradiated targets were allowed to cool for a period of time. After cooling, the irradiated samples were dissolved in double distilled water. A stock solution of 1000 mg L$^{-1}$ Cs (I) and Co (II) ions was prepared separately by dissolving the essential amounts of Cesium chloride (CsCl) and cobalt chloride (CoCl$_2$·6H$_2$O) in a certain volume of double distilled water, whereby it has become possible to prepare the desired concentrations of the adsorbate solution by suitable dilution.

**Characterization**

The physical and chemical properties of the soil samples, in the present study, were characterized using X-Ray Fluorescence (XRF) analysis using on a PANalytical PW2404 X-ray fluorescence spectrometer for determining the chemical composition. The functional groups of soil samples were determined using FT–IR spectrophotometer of type Nicolet 10 from Thermo (USA). The phase structure was investigated using X-Ray diffractometer. The analysis was achieved using X-Ray diffractometer. The analysis was achieved using PANalytical X’Pert PRO through monochromator, Cu-radiation ($\lambda$=1.5418 Å) run at a voltage 45 kV, current 35 mA and scanning speed 0.03°.sec$^{-1}$.

**Batch experiments**

Sorption studies of Cs (I) and Co (II) ions onto soil samples were performed by stirring a definite weight of soil samples with 10 mL of 100 mg.L$^{-1}$
metal ion solution of Cs (I) and Co (II) ions, in closed glass bottles. The solutions were traced with the corresponding radioactive isotopes $^{137}$Cs and $^{60}$Co and their initial pH were adjusted at 6 and 5.8 values by addition of 0.1 M HCl or 0.1 M NaOH solutions. At certain time intervals, the supernatants were separated by centrifugation and subjected to radiometric analysis to determine the activity of the studied radionuclides. The radioactivity of studied isotopes was $\gamma$-assayed using a multi-channel counter connected to a NaI scintillation detector. The uptake percentage (U%) was calculated using the following equation:

$$U\% = \frac{A_f - A_i}{A_i} \times 100$$ (1)

where: $A_i$ and $A_f$ are the initial and final radioactivity levels of the studied radionuclides. The maximum sorption capacity ($Q_{\text{max}}$, mg.g$^{-1}$) for the two soil samples was determined by repeated equilibration of certain weights of soil samples with a definite volume of 100 mg.L$^{-1}$ Cs(I) and Co(II) solutions spiked with the corresponding radioactive isotopes at certain initial pH values. The samples were stirred at room temperature for 24 h. Then, the supernatants were drag, centrifuged and $\gamma$-radioassayed. An equal volume of the original solution of studied ions was added to the soil samples and stirred till achieving equilibrium. The former procedure were repeated till no additional sorption occurred for both Cs(I) and Co(II). The sorbed amount ($q_e$) onto soil sample for each equilibration process was determined from the following equation [15]:

$$q_e = (C_0 - C_e) \times \frac{V}{m}$$ (2)

where: $C_0$ and $C_f$ are the initial and final concentrations (mg.L$^{-1}$) of Cs(I) and Co(II) metal ions, V is the volume of solution (L) and m is the weight of soil sample (g). The effect of pH value was studied at ranges from 1 to 11 for Cs(I) and Co(II) solutions. After equilibration, the supernatants radioactivity was radiometrically an additional set of experiments were performed as previously described to clarify the effect of certain parameters.

Results and Discussion
The specific surface area had been determined according to Brunauer-Emmett-Teller (BET). The soil samples have surface area of 10.532 and 17.970 m$^2$.g$^{-1}$ for DNS/C and DNS/Q respectively.

Characterization
The chemical composition of DNS/C and DNS/Q soil samples were determined using quantitative XRF analysis and results are presented in Table (1). The XRF analysis shows that calcium oxide (CaO) was the most abundant component (43.83%) in DNS/C sample while silica (SiO$_2$) was the most abundant component (39.65%) in DNS/Q sample. The oxides silica (SiO$_2$), sodium oxide (Na$_2$O) and magnesium oxide (MgO) were detected in low percentages amounting to 5.3, 2.53 and 2.1 % in DNS/C sample. Also, other oxides such as titanium oxide (TiO$_2$), alumina (Al$_2$O$_3$), iron oxide (Fe$_2$O$_3$), manganese oxide (MnO), potassium oxide (K$_2$O), phosphorus pentoxide (P$_2$O$_5$) and sulfur trioxide (SO$_3$) were detected at levels less than 2 % of the total weight of DNS/C soil sample. On other hand, the DNS/Q sample was mainly composed of calcium oxide (CaO), alumina (Al$_2$O$_3$), magnesium oxide (MgO) and iron oxide (Fe$_2$O$_3$) with percentages amounting to 22.36, 8.26, 3.50 and 2.53 %, respectively. Other oxides like potassium oxide (K$_2$O), titanium oxide (TiO$_2$), manganese oxide (MnO), sodium oxide (Na$_2$O), phosphorus pentoxide (P$_2$O$_5$) and sulfur trioxide (SO$_3$) were detected at levels less than 2 % of the total composition.

The loss on the ignition of DNS/C and DNS/Q soil samples was found to be 35.06 and 18.16 %. The high loss, due to ignition, could be mainly attributed to calcium carbonate decomposition that lead to formation of CaO and carbon dioxide (CO$_2$) [16]. The FT-IR spectrum of DNS/C and DNS/Q are shown in Figure (2). The bands detected at ~1485 ~1080, ~1788, ~855, and ~711 cm$^{-1}$ could be referred to absorption vibration of CO$_2$ groups in both soil samples. They were reported as common features of carbonate ions in calcium carbonate and could be considered as the basic patterns of vibration for this molecule [17, 18].

The frequency at 1788 cm$^{-1}$ could be ascribed to C=O stretching vibration. The frequency at 2522 cm$^{-1}$ was identified for calcite in the samples [19, 20]. The absorption bands at 2920–2854 cm$^{-1}$ was attributed to C–H vibration [21]. The band at 2626 cm$^{-1}$ is a shoulder band accompanying absorption of carbonate while that observed at 2522 cm$^{-1}$ was due to calcite/dolomite discrimination [22]. The broad range of 3600–3100 cm$^{-1}$ corresponded to O–H bonding stretching vibrational of water of hydration. The band exhibited at 1670–1600 cm$^{-1}$

could be referred to O-H bending vibrational modes for DNS/Q soil sample. The frequency at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 455 and 540 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23]. The band observed at 3686 cm$^{-1}$ could be attributed to metal oxygen stretching vibrational modes [23].

The crystalline structure for both soil samples was analyzed using X-ray diffraction technique and the data obtained are given in Figure (3). DNS/C exhibited characteristics peaks of calcite, magnesium at 2θ values of 23.29°, 29.62°, 36.24°, 39.66°, 43.39° and 47.76°, which correlated with (hkl) indices of (012), (104), (110), and (113), (202), and (024), respectively. The data illustrate that the crystal structure of the soil was trigonal (hexagonal axes) [25]. The strongest peak for DNS/C sample was observed at 2θ value of 29.62° with intensity 100% and d-spacing 3.02. However, peaks were also observed at 21.04°, 26.80°, 46.00°, 50.33°, 26.39°, 27.41°, and 33.28° that could be ascribed to the minerals quartz, graphite, halite, and hematite, respectively [26-29]. X-ray diffraction pattern of DNS/Q exhibited characteristics peaks of quartz at 2θ values of 21.14°, 26.89°, 36.83°, 39.79°, 40.59°, 42.72°, 46.08°, 50.41° and 55.24° that correlated with (hkl) indices of (100), (101), (110), and (102), (111), (200), (201), (112) and (103) respectively. The data illustrate that DNS/Q soil sample had a trigonal crystal structure (hexagonal axes) [30].

Table (1): Chemical composition of DNS/C and DNS/Q soil samples

<table>
<thead>
<tr>
<th>Compound</th>
<th>Percentage (% w/w)</th>
<th>DNS/C</th>
<th>DNS/Q</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica(SiO$_2$)</td>
<td>5.3</td>
<td>39.65</td>
<td></td>
</tr>
<tr>
<td>Titanium oxide (TiO$_2$)</td>
<td>0.12</td>
<td>0.46</td>
<td></td>
</tr>
<tr>
<td>Alumina (Al$_2$O$_3$)</td>
<td>0.88</td>
<td>8.26</td>
<td></td>
</tr>
<tr>
<td>Iron oxide (Fe$_2$O$_3$)</td>
<td>1.08</td>
<td>2.53</td>
<td></td>
</tr>
<tr>
<td>Manganese oxide (MnO)</td>
<td>0.04</td>
<td>0.05</td>
<td></td>
</tr>
<tr>
<td>Magnesium oxide (MgO)</td>
<td>2.1</td>
<td>3.50</td>
<td></td>
</tr>
<tr>
<td>Calcium oxide (CaO)</td>
<td>43.83</td>
<td>22.36</td>
<td></td>
</tr>
<tr>
<td>Sodium oxide (Na$_2$O)</td>
<td>2.53</td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td>Potassium oxide (K$_2$O)</td>
<td>0.32</td>
<td>1.98</td>
<td></td>
</tr>
<tr>
<td>Phosphorus pentoxide (P$_2$O$_5$)</td>
<td>&lt; 0.01</td>
<td>&lt; 0.01</td>
<td></td>
</tr>
<tr>
<td>Chlorine(Cl)</td>
<td>7.89</td>
<td>2.06</td>
<td></td>
</tr>
<tr>
<td>Sulfur trioxide (SO$_3$)</td>
<td>0.15</td>
<td>0.09</td>
<td></td>
</tr>
</tbody>
</table>

Fig. (2): FT-IR spectra of: a) DNS/C and b) DNS/Q soil samples

Fig. (3): XRD patterns of soil samples: a) DNS/C and b) DNS/Q

The strongest peak for DNS/Q was observed at a 2θ value of 26.89° with 100% intensity and d-spacing 3.22. There are peaks observed also at 29.73°, 43.52°, 47.90°, 48.87° and 31.07° and 41.34°, 27.72 and 8.75° corresponding to calcite-magnesium, halite, dolomite, sylvine and muscovite-2M1 minerals respectively [25, 31-34].

The crystallite size (t) of the soil samples was determined using data of XRD by applying Debye-Scherrer equation:

$$t = \frac{K \cdot \lambda}{\beta \cdot \cos \theta}$$

(3)

where: $t$ is the crystallite soil sample size in (nm), $k$ is a constant relying on crystallite shape and has the value of 0.89, $\lambda$ is the wavelength of X-rays and has the value of 0.1542 nm, $\theta$ is the angle of diffraction given in (degree) and $\beta$ is the peak width of full diffraction at half maximum intensity (FWHM). The mean crystallite sizes, are estimated using Scherrer equation, were 41.34 and 100.17 nm for DNS/C and DNS/Q respectively. The values demonstrate the nanosized structures of the applied soil samples.

**Sorption study**

*Effect of time*

The change in uptake percent of Cs(I) and Co(II) with time was recorded at different periods ranging from 5 min to 48 h. The sorption was performed using batch technique and obtained data are shown in Figure (4). The data illustrate that the uptake of Cs(I) and Co(II) by DNS/C and DNS/Q soil samples had two different steps. The first step included a sharp increase in the uptake rate with contact time up to 4 h and the final step referred to a slow uptake rate till reaching equilibrium after 24 h. The uptake percentages of Cs(I) were 22.59 and 47.14 % for sorption onto DNS/C and DNS/Q soil samples while the values 79.20 and 90.15 % were recorded for sorption of Co(II) onto DNS/C and DNS/Q samples, respectively. The rapid sorption of Cs(I) and Co(II) by soil samples can illustrate that the increasing of active sites at initial step available for sorption of studied ions related to the concentration of Cs(I) and Co(II) ions interpreting the increasing of sorption rate.

By increasing sorption, active sites on soil surface started to decrease and ions started to compete between themselves for active sites. Subsequently, the uptake percent went down [35]. The maximum sorption amounts ($Q_{\text{max}}$) were determined and found to be 1.16 and 2.39 mg.g$^{-1}$ for sorption of Cs (I) onto DNS/C and DNS/Q while $Q_{\text{max}}$ for sorption of Co(II) onto DNS/C and DNS/Q were 4.28 and 4.76 mg.g$^{-1}$, respectively.

![Fig. (4): Effect of contact time on uptake of: a) cesium and b) cobalt ions onto the studied soil samples (pH$_{i(Ca)}$ = 6 ; pH$_{i(Co)}$ = 5.8 ; V/m = 100 mL.g$^{-1}$; $c_{o}=20$ mg.L$^{-1}$ ; temp. = 25 °C).](image)

### 3.2.2. Effect of pH

Solution pH is an important dominant parameter in sorption processes because of its effect on the surface properties of soil samples and the ionic form of the metal ion in aqueous solution. [36]. The effect of pH value on the uptake percentages of soil samples for Cs(I) and Co(II) ions is illustrated in Figure (5). The uptake percentages of cesium ion were increased by increasing the pH value from 2 to 7. This is because $H_3O^+$ is of a higher concentration at lower pH values and competed with cesium ions for active sites [37, 38]. With increasing pH, the concentration of $H_3O^+$ decreased and the deprotonation of active sites on soil surface increased that enable these sites to attract cesium ions and hence the uptake of cesium increased. At higher pH values, the uptake of Cs(I) was slightly decreased with increasing pH value up to 11. A similar behavior was reported by other investigators [39]. The maximum sorption of cesium was detected at pH 6.
The uptake of Co(II) on the applied soil samples is shown in Figure (5). The plots clarify that the uptake of Co(II) increased with rising pH value from 1 up to 6. At higher pH values, the Co(II) uptake slightly increased till reaching equilibrium, where the uptake percentage attained a constant value with rising pH value up to 11. To verify the solubility extent of both metal ions under the applied experimental conditions, a blank solution of metal ions had been studied without presence of soil samples to achieve the solubility range of these metal ions. The results show that Cs(I) ions are completely soluble over the studied pH range extending from 1-11, where Cs⁺ is the predominant species over this pH range. On the other hand, Co(II) ions exhibited different solubilities over different pH values, Co(II) precipitation started at pH ~ 6.5 at which the removal was attributed to the precipitation of Co(II) ions. At these conditions, Co(OH)₂ were the predominant species [40]. The uptake percent (U%) of Cs(I) ions onto DNS/C and DNS/Q samples attained the values 23.2 and 47.8 % at initial pH 6 while Co(II) ions attained the values 78.9 and 90.1 % at initial pH 5.8, respectively. Based on these data, pH 6 and 5.8 was applied in the rest of experiments for studying the sorption of Cs(I) and Co(II) ions on the applied soil samples.

**Effect of competing ion**

The influence of the presence of varied initial concentrations of different cations as Na⁺ (univalent), Ca²⁺, Mg²⁺ and Mn²⁺ (bivalent) and Al³⁺ (trivalent) on uptake of Cs(I) and Co(II) onto the used soil samples is clarified in Figure (6). The data illustrate that the uptake of Cs(I) decreased with presence of increased concentration of the metal ions Na⁺, Ca²⁺, Mg²⁺, Mn²⁺ and Al³⁺. The decrease in Cs(I) uptake increased with increasing the initial concentration of these competing ions up to 0.4 M, Figure (6a). The closer inspection to these plots clarifies a sharp decrease in Cs(I) uptake with rising the initial concentration of all competing cations up to 0.1 M. At higher concentration, the uptake percent of Cs(I) slightly decreased with increasing the initial concentration of all competing ions up to 0.4 M. The deteruous effect of the applied competing cations is on the order: Al³⁺ > Mg²⁺ > Mn²⁺ > Ca²⁺ > Na⁺. Additionally, the role of the previously mentioned interfering cations on the uptake of Co(II) ions is illustrated in Figure (6b). The plots exhibit a decrease in the uptake of Co(II) with the presence of such cations and this depress in uptake percent of Co(II) is directly proportional to the initial concentration of the applied competing cations.

**Sorption kinetics**

The sorption rate of Cs(I) and Co(II) onto the soil samples was illustrated using pseudo-first-order and pseudo-second-order models.

**Pseudo-first-order kinetic model**

The pseudo-first-order model was proposed in sorption of liquid to solid system that depends on the solid concentration. It was given by the following equation [41]:

$$\log(q_e - q_t) = \log q_e - \frac{K_f t}{2.303}$$

(3)
INVESTIGATING THE SORPTION …..

Fig. (6): Effect of competing ions on uptake of: a) Cs(I) and b) Co(II) ions onto DNS/C soil sample (pH_{i(Cs)} = 6; pH_{i(Co)} = 5.8; t_{eq} = 24h; V/m = 100 mL.g^{-1}; C_{0} = 20 mg.L^{-1}; temp. = 25^\circ C)

where: q_{e}, q_{t} are the adsorbed amount of Cs(I) or Co(II) onto soil sample (mg.g^{-1}) at equilibrium and at time t and k_{i} is the rate constant of pseudo-first-order (min^{-1}). The relation of log (q_{e} - q_{t}) and time for Cs(I) and Co(II) sorption onto studied soil samples is given in Figures (7). The model constants were determined from the slope and intercept of the straight lines of the corresponding plots and their values along with the correlation coefficients (R^2) are illustrated in Table (2). This data illustrate that the adsorbed amounts (q_{e}) of Cs(I) on DNS/C and DNS/Q soil samples, after reaching equilibrium, were found to be 0.28 and 0.43 mg.g^{-1}, respectively while the experimental values were 0.45 and 0.94 mg.g^{-1}. The data indicate that q_{e} values estimated using pseudo-first-order kinetic model weren’t consistent with the experimental q_{e} values for both Cs(I) and Co(II). This confirms that the kinetics of Cs(I) and Co(II) sorption using the soil samples did not follow the pseudo-first-order reaction. In order to find a more reliable description of the sorption kinetics, a pseudo-second-order kinetic equation was employed.

samples were found to be 0.91 and 1.00 mg.g^{-1}, respectively while the experimental values were 1.58 and 1.79 mg.g^{-1}. The data indicate that q_{e} values estimated using pseudo-first-order kinetic model weren’t consistent with the experimental q_{e} values for both Cs(I) and Co(II). This confirms that the kinetics of Cs(I) and Co(II) sorption using the soil samples did not follow the pseudo-first-order reaction. In order to find a more reliable description of the sorption kinetics, a pseudo-second-order kinetic equation was employed.

Fig. (7): Plots of pseudo-first-order for sorption of: a) Cs(I) and b) Co(II) ions onto the studied soil samples

Table (2): Pseudo-first-order constants for sorption of Cs(I) and Co(II) ions onto the studied soil samples
Pseudo-second-order kinetic model

Pseudo-second-order model was used to describe the chemisorption kinetic of sorbate ions onto sorbent surface. The mathematical linear form of this model is given below [42]:

\[
\frac{t}{q_t} = \frac{1}{k_s q_e^2} + \frac{1}{q_e} t
\]

(4)

where: \(k_s\) is the pseudo-second-order rate constant (g mg\(^{-1}\)min\(^{-1}\)), the product (\(k_s q_e^2\)) is the initial sorption rate represented as \(h\). The relation between \(t/q_t\) and \(t\) for the sorption of Cs(I) or Co(II) ions onto the studied soil samples is shown in Figure (8). The constant of pseudo-second-order model were estimated from the slope and intercept of the straight lines of the corresponding plots. The values of these constants and the correlation coefficients (R\(^2\)) are given in Table (3).

The data illustrate that adsorbed amounts (\(q_e\)) of Cs(I) on DNS/C and DNS/Q soil samples after reaching equilibrium were found to be 0.43 and 0.91 mg g\(^{-1}\), while the experimental values were 0.45 and 0.94 mg g\(^{-1}\). The adsorbed amounts (\(q_e\)) of Co(II) on DNS/C and DNS/Q soil samples were found to be 1.62 and 1.81 mg g\(^{-1}\), respectively, while the experimental values were 1.58 and 1.79 mg g\(^{-1}\). The results illustrate that adsorbed amounts values determined from pseudo-second-order model were consistent with the experimental \(q_e\) values for sorption of both Cs(I) and Co(II) ions and the correlation coefficient (R\(^2\)) is extremely high and closer to unity (R\(^2\) > 0.998). This illustrated that sorption of Cs(I) and Co(II) onto the surface of applied soil samples could be represented by the kinetic of pseudo-second-order kinetics. Hence, the rate-determining step in the sorption process is a chemisorption process dependent on the number of active sites on soil surface and the initial concentration of metal ions [43].

**Table (3): Pseudo-second-order constants for sorption of Cs(I) and Co(II) ions onto the studied soil samples**

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<thead>
<tr>
<th>Cations</th>
<th>Model constants</th>
<th>Soil samples</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>DNS/C</td>
</tr>
<tr>
<td>Cs(I)</td>
<td>(k_s) (min(^{-1}))</td>
<td>0.050231</td>
</tr>
<tr>
<td></td>
<td>(q_e) (mg g(^{-1}))</td>
<td>0.430674</td>
</tr>
<tr>
<td></td>
<td>(h) (mg g(^{-1}) min(^{-1}))</td>
<td>0.009318</td>
</tr>
<tr>
<td></td>
<td>R(^2)</td>
<td>0.99899</td>
</tr>
<tr>
<td></td>
<td>S.D</td>
<td>2.61E-02</td>
</tr>
<tr>
<td>Co(II)</td>
<td>(k_s) (mg g(^{-1}) min(^{-1}))</td>
<td>0.013756</td>
</tr>
<tr>
<td></td>
<td>(q_e) (mg g(^{-1}))</td>
<td>1.628638</td>
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<tr>
<td></td>
<td>(h) (mg g(^{-1}) min(^{-1}))</td>
<td>0.036487</td>
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<tr>
<td></td>
<td>R(^2)</td>
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<td></td>
<td>S.D</td>
<td>6.58E-03</td>
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</table>
Sorption thermodynamics
To investigate the effect of temperature on the sorption of $^{134}$Cs and $^{60}$Co radionuclides onto the employed soil samples, the sorption experiments were conducted at different temperatures having the values 30, 35, 40 and 45 °C and the data obtained are given in Figures (9). The plots show that the increase in temperature had a negative effect on the sorption of cesium radionuclide where the sorption slightly decreased with increasing temperature Figure (9-a). This result was similar to that reported by Tertre and Wu [44]. This data indicated that sorption of cesium radionuclides onto the clay surface was favored at lower temperatures. On the other hand, the sorption of cobalt was increased with rising temperature from 30 °C to 45 °C, suggesting that sorption of cobalt onto soil samples is promoted at a higher temperature.

Fig. (9): Effect of temperature on uptake of: a) Cs(I) and b) Co(II) ions onto the studied soil samples ($C_0 = 20$ mg L$^{-1}$; pH$_{0,\text{Cs}} = 6$; pH$_{0,\text{Co}} = 5.8$; V/m = 100 mL.g$^{-1}$; $t_{eq} = 24$ h; temp. = 25 °C)

The revealed sorption data were used to estimate the thermodynamic parameters. The entropy change ($\Delta S^o$) and enthalpy change ($\Delta H^o$) were determined from Van’t Hoff equation [45]:

$$\ln K_d = \frac{\Delta S^o}{R} - \frac{\Delta H^o}{RT}$$

where: $\Delta S^o$ is the entropy change (kJ.mol$^{-1}$.K$^{-1}$), $\Delta H^o$ is the change of enthalpy (kJ.mol$^{-1}$), $R$ is the ideal gas constant (8.315×10$^{-3}$ kJ.mol$^{-1}$.K$^{-1}$) and $T$ is the absolute temperature (°K). The distribution coefficient ($K_d$) for sorption of Cs(I) and Co(II) was determined using the equation:

$$K_d = \frac{C_a-C_e}{C_e} \times \frac{V}{m}$$

where: $K_d$ is the distribution coefficient (mL.g$^{-1}$), $C_a$ and $C_e$ are the initial and equilibrium concentrations of Cs(I) and Co(II) ions in solution (mg.L$^{-1}$) respectively, $V$ is the volume of solution (mL) and $m$ is the sorbent mass (g). The relation of $\ln K_d$ versus $1/T$ for sorption of Cs(I) and Co(II) onto the soil samples is shown in Figure (10). $\Delta H^o$ and $\Delta S^o$ values were calculated from the slope and the intercept of the straight lines of the corresponding plots and the values are listed in Tables (4). The free energy change ($\Delta G^o$), is determined using the following equation

$$\Delta G^o = -RT \ln K_d$$

where: $\Delta G^o$ is the change of Gibbs free energy in (kJ.mol$^{-1}$). From Eq. (5), Eq. (7) can be written in the form:

$$\Delta G^o = \Delta H^o - T \Delta S^o$$

Sorption of cesium radionuclide onto the studied clays gives negative $\Delta H^o$ values, which indicate the exothermic nature of this sorption process at the studied experimental conditions. The positive values of $\Delta H^o$ for cobalt radionuclide indicate that the sorption process of this radionuclide onto the studied soils is an endothermic process. The negative values of $\Delta S^o$ recorded for Cs (I) ions onto the studied soils indicate less randomness at the soil-solution interface during sorption processes. The positive values of $\Delta S^o$ recorded for sorption Co(II) ions onto studied soils indicate the increased randomness at the soil-solution interface during the sorption processes. For $\Delta G^o$, the positive values of $\Delta G^o$ for cesium radionuclide with increasing temperature show that its sorption onto studied soils is more thermodynamically feasible at lower temperatures and is non-spontaneous. On the contrary, its negative values obtained under our experimental conditions for
Co(II) ions indicate that the present sorption processes of Co(II) ions are thermodynamically feasible and spontaneous in nature. The increase in the values of \(-\Delta G^o\) with increasing temperature from 30 °C to 45 °C demonstrates that the sorption processes of cobalt radionuclide onto soil samples, used in the present work, are more feasible at higher temperatures.

![Graph](image1)

**Fig. (10):** Plots of ln Kd versus 1/T for sorption of: a) Cs(I) and b) Co(II) ions onto the studied soil samples.

**Table (4):** The values of thermodynamic parameters for sorption process of Cs(I) and Co(II) the studied soil samples

<table>
<thead>
<tr>
<th>Soil Sample</th>
<th>Temp. (K)</th>
<th>Element</th>
<th>(\Delta G^o) (kJ mol(^{-1}))</th>
<th>(\Delta H^o) (kJ mol(^{-1}))</th>
<th>(\Delta S^o) (J K(^{-1}) mol(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNS/C</td>
<td>303</td>
<td>Cs(^+)</td>
<td>-0.329</td>
<td>0.699</td>
<td>0.003150</td>
</tr>
<tr>
<td></td>
<td>308</td>
<td></td>
<td></td>
<td></td>
<td>0.07390</td>
</tr>
<tr>
<td></td>
<td>313</td>
<td></td>
<td></td>
<td></td>
<td>0.0774</td>
</tr>
<tr>
<td></td>
<td>318</td>
<td></td>
<td></td>
<td></td>
<td>0.08025</td>
</tr>
<tr>
<td>DNS/Q</td>
<td>303</td>
<td>Co(^{2+})</td>
<td>0.069</td>
<td>1.068</td>
<td>0.003150</td>
</tr>
<tr>
<td></td>
<td>308</td>
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<td>318</td>
<td></td>
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<td>0.08025</td>
</tr>
</tbody>
</table>

**Conclusion**

In the current study, sorption of Cs(I) and Co(II) ions onto DNS/C and DNS/Q soil samples collected from El-Dabaa site, Egypt, was investigated. DNS/Q have a high affinity for sorption of Cs(I) and Co(II) ions under optimal conditions, whereas DNS/C have a lower affinity for sorption for both metal ions as specific surface area are 10.532 and 17.970 m\(^2\) g\(^{-1}\), d-spacing results from XRD are 3.02 and 3.32 nm for DNS/C and DNS/Q respectively and XRF analysis illustrates that the percentages of Alumina (Al\(_2\)O\(_3\)) for DNS/C and DNS/Q are 8.8 and 8.26% form the total weight that have a great effect on the uptake percentage discussed in competing ion effect. Kinetic studies revealed that sorption of El-Dabaa soil soils for Cs(I) and Co(II) radionuclides were compatible with pseudo-second-order model estimated that the sorption process followed it and has a chemisorption behaviour.

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**References**

8. M. A. Barakat, R. Kumar, Synthesis and characterization of porous magnetic silica composite for the removal of heavy metals.


