Modelling Dispersion Influences on Chromium Transport in Heterogeneous Silty and Fine Sand Formation, Port Harcourt Industrial Lay-Out, Rivers State of Nigeria

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Abstract: This paper examined chromium transport through dispersion and storage coefficient influences. The study has monitored the deposition of chromium in silty and fine sand formation. The developed model expressed the behaviour of chromium in the study location, storage coefficient and dispersions of chromium were examined through the behaviour of chromium transport in silty and fine sand formation, the behaviour were expressed through graphical representation, the fluctuation on concentration reflect the influences from porosity variation thus dispersion and storage coefficient. This generated slight accumulation of chromium deposition in silty and fine sand formation, this condition was examined through the rate of chromium deposition; some fluctuations were experienced from the experimental values for model validation. The heterogeneous settings in the formation were also observed, the developed model was compared with other experimental values, and both parameters expressed favourable fits validating the model.

Keywords: Dispersion, Storage Coefficients, Chromium and Heterogeneous Formation

1. Introduction

For several years experts in different dimension has been developed various sources of heavy metal, more soil were examined to deposited over a billion individual cells and estimation of 104–105 distinct genomes per gram of soil [15, 18, 19 and 23], it is observed that bacteria in soil are the reservoirs for much of Earth’s genetic biodiversity. This vast phylogenetic and efficient diversity can be attributed to the dynamic physical and chemical heterogeneity of soil, the results has express spatial and temporal separation of microorganisms including chemical constituents [8-10]. More so it is observed from high diversity of carbon (C) – rich compounds in soils, it has been observed that the ability of each taxon to compete for only a subset of resources could also contribute to the high diversity of bacteria including chemicals of natural origin in soils through resource partitioning [9-11, and 13]. Indeed, [10, 14 and 17] this were thoroughly express through distinct substrate preferences by broad microbial groups in grassland soils relating it to natural chemical deposition and C resource partitioning it, such condition demonstrate some key that contribute patterning of bacterial co-existence in model communities on plant surfaces [7, 12-13, and 14]. The development of high-throughput tools to evaluate the composition of soil bacterial communities is rapidly contributing to an enhanced perceptive of bacterial diversity and biogeographically distribution [4-6, and 11]. However, our ability to evaluate the purpose of different bacterial taxa has not reserved pace [15 - 16]. There are some limits in defining the consequences shifting community in response
to environmental change [1-3, and 16]. Numerous concepts applied to monitor the trace of the bacteria for this reason, the application of tracer molecules such as stable-isotopes and the thymidine analog, 3-bromodeoxyuridine (BrdU), have been widely accepted. Effort to connect phylogeny to function will be easier through this process. Stable-isotopes, particularly the heavy carbon isotope 13C, have been regularly applied to recognize microbial community members capable of catabolizing particular substrates [17-21, and 23]. This method requires separation of nucleic acids based on buoyant density, so high concentrations of isotopically labeled substrate are needed. This approach was first applied to the study of bacterial populations over a decade ago [21-22] and it has since been used to categorize soil bacterial taxa that respond to various environmental stimuli [2-3 and 6]. Recently, BrdU incorporation has been shown to detect a broad diversity of bacterial phyla in marine systems [4-8 and 19] [and fungal taxa in temperate [13, 22] and boreal forest soils [5-8 and 20].

2. Theoretical Background

The behaviour of chromium in the soil has been observed by experts in different dimension, the rate of migration in soil and water environment are monitored through various way under the influences of flow in the soil through various formation characteristics, the study of storage coefficient including dispersion of chromium in Phreatic bed influencing it transport process has not been thoroughly expressed. This implies that the migration of the chromium through these sources has not been evaluated from this study, the rate of dispersions of the chromium are through these flows, these are base on the fluctuation rate on its permeation of the formation between the lithology in the study area.

3. Governing Equation

\[
D \frac{d^2c}{dx^2} - ST \frac{dc}{dx} + V \frac{dc}{dx} = 0
\]  
(1)

\[
D \frac{d^2c}{dx^2} - (ST - V) \frac{dc}{dx} = 0
\]  
(2)

Let \( C = \sum_{n=0}^{\infty} a_n x^n \)

\[
C^0 = \sum_{n=0}^{\infty} n a_n x^{n-1}
\]

\[
C^{-1} = \sum_{n=2}^{\infty} n(n-1) a_n x^{n-2}
\]

\[
D \sum_{n=2}^{\infty} n(n-1) a_n x^{n-2} - (ST - V) \sum_{n=0}^{\infty} n a_n x^{n-1} = 0
\]  
(3)

Replace \( n \) in the 1st term by \( n+2 \) and in the 2nd term by \( n+1 \), so that we have;

\[
D \sum_{n=2}^{\infty} (n+2)(n+1) a_{n+2} x^n - (ST - V) \sum_{n=0}^{\infty} (n+1) a_{n+1} x^n = 0
\]  
(4)

i.e. \( D(n+2)(n+1) a_{n+2} - (ST - V)(n+1) a_{n+1} = 0 \)

\[
a_{n+2} = \frac{(ST - V)(n+1) a_{n+1}}{D(n+2)(n+1)}
\]  
(5)

\[
a_{n+2} = \frac{(ST - V) a_{n+1}}{D(n+2)}
\]  
(6)

\[
\text{for } n = 0, \ a_2 = \frac{(ST - V) a_1}{2D}
\]  
(7)
for \( n = 1, a_i = \frac{(ST - V_i)a_i}{3D} = \frac{(ST - V_i)^2 a_i}{2D \bullet 3D} \) \hspace{1cm} (9)

for \( n = 2, a_i = \frac{(ST - V_i)a_i}{4D} = \frac{(ST - V_i) \bullet (ST - V_i)a_i}{3D \bullet 2D} = \frac{(ST - V_i)^3 a_i}{4D \bullet 3D \bullet 2D} \) \hspace{1cm} (10)

for \( n = 3, a_i = \frac{(ST - V_i)a_i}{5D} = \frac{(ST - V_i)}{5D \bullet 4D \bullet 3D \bullet 2D} \) \hspace{1cm} (11)

for \( n; a_n = \frac{(ST - V_i)^{n-1} a_i}{D^{n-1} n!} \) \hspace{1cm} (12)

\[
C(x) = a_0 + a_1 x + a_2 x^2 + a_3 x^3 + a_4 x^4 + a_5 x^5 + \ldots + a_n x^n
\]

\[
C(x) = a_0 + a_1 \frac{(ST - V_i)a_i}{2D} + a_2 \frac{(ST - V_i)a_i}{3D^2} + a_3 \frac{(ST - V_i)a_i}{4D^3} + a_4 \frac{(ST - V_i)a_i}{5D^4} + \ldots
\]

\[
C(x) = a_0 + a_1 \left[ x + \frac{(ST - V_i)x^2}{2!D} + \frac{(ST - V_i)x^3}{3!D^2} + \frac{(ST - V_i)x^4}{4!D^3} + \frac{(ST - V_i)x^5}{5!D^4} + \ldots \right]
\]

\[
C(x) = a_0 + a_1 \ell \frac{(ST - V_i)}{D}
\]

Subject equation (16) for the following boundary conditions

\[
C(o) = 0 \text{ and } C(o) = H
\]

\[
C(x) = a_0 + a_1 \ell \frac{(ST - V_i)}{D}
\]

\[
C(o) = a_0 + a_1 = 0
\]

i.e. \( a_0 + a_1 = 0 \)

\[
C^l(x) = \frac{(ST - V_i)}{2!D} a_1 \ell \frac{(ST - V_i)}{D}
\]

\[
C^l(o) = \frac{(ST - V_i)}{2!D} a_1 = H
\]

\[
a_1 = \frac{HD}{ST - V_i}
\]

(18)

Substitute (18) into equation (17)

\[
a_1 = a_0
\]

\[
\Rightarrow a_0 = -\frac{HD}{ST - V_i}
\]

(19)

Hence the particular solution of equation (16) is of the form:

\[
C(x) = -\frac{HD}{ST - V_i} + \ell \frac{(ST - V_i)}{D}
\]
4. Materials and Method

Standard laboratory experiment where performed to monitor the concentration of chromium depositions indifferent formation, applying AAS to monitor various rates of concentration in different strata, the soil strata were collected in sequences base on the structural deposition at different locations, this samples collected at different location generated variation at different depth producing different migration of chromium concentration through pressure flow at different strata, the experimental result are applied to be compared with the theoretical values to determined the validation of the model.

5. Result and Discussion

Results and discussion are presented in tables including graphical representation of chromium concentration.

\[
C(x) = \frac{HD}{ST - V_i} \left[ \ell \left( \frac{ST - V_i}{L} \right)^{-1} \right]
\]  

(20)

*Figure 1. Concentration of chromium at Different Depth.*

*Figure 2. Predicted and Validate Concentration of chromium at Different Depth.*
Figure 3. Concentration of chromium at Different Depth.

Figure 4. Predicted and Validate Concentration of chromium at Different Depth.

Figure 5. Concentration of chromium at Different Depth.
Figure 6. Predicted and Validate Concentration of chromium at Different Depth.

Figure 7. Concentration of chromium at Different Depth.

Figure 8. Predicted and Validate Concentration of chromium at Different Depth.
Table 1. Concentration of chromium at Different Depth.

<table>
<thead>
<tr>
<th>Depth [M]</th>
<th>chromium Concentration</th>
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</thead>
<tbody>
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<td>3</td>
<td>1.13E-05</td>
</tr>
<tr>
<td>6</td>
<td>2.20E-05</td>
</tr>
<tr>
<td>9</td>
<td>3.31E-05</td>
</tr>
<tr>
<td>12</td>
<td>4.41E-06</td>
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<td>39</td>
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Table 2. Predicted and Validate Concentration of chromium at Different Depth.

<table>
<thead>
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<th>Depth [M]</th>
<th>Predictive Values</th>
<th>Experimental Values</th>
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<tbody>
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<tr>
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<td>36</td>
<td>1.32E-08</td>
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<td>39</td>
<td>1.43E-08</td>
<td>1.52E-08</td>
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Table 3. Concentration of chromium at Different Depth.

<table>
<thead>
<tr>
<th>Time Per Day</th>
<th>chromium Concentration</th>
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<tbody>
<tr>
<td>10</td>
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<td>20</td>
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<td>90</td>
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<td>120</td>
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<td>130</td>
<td>2.00E-02</td>
</tr>
<tr>
<td>140</td>
<td>2.16E-02</td>
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Table 4. Predicted and Validate Concentration of chromium at Different Depth.

<table>
<thead>
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<th>Predictive Values</th>
<th>Experimental Values</th>
</tr>
</thead>
<tbody>
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<td>3.00E-03</td>
<td>3.04E-03</td>
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<tr>
<td>30</td>
<td>4.41E-03</td>
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<td>6.18E-03</td>
<td>6.09E-03</td>
</tr>
<tr>
<td>50</td>
<td>7.72E-03</td>
<td>7.54E-03</td>
</tr>
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</table>

Table 5. Concentration of chromium at Different Depth.

<table>
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<tr>
<th>Depth [M]</th>
<th>chromium Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>2.62E-05</td>
</tr>
<tr>
<td>6</td>
<td>5.25E-05</td>
</tr>
<tr>
<td>9</td>
<td>7.88E-05</td>
</tr>
<tr>
<td>12</td>
<td>1.05E-05</td>
</tr>
<tr>
<td>15</td>
<td>1.31E-04</td>
</tr>
<tr>
<td>18</td>
<td>1.51E-04</td>
</tr>
<tr>
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<td>2.36E-06</td>
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<td>3.41E-06</td>
</tr>
<tr>
<td>42</td>
<td>3.67E-06</td>
</tr>
</tbody>
</table>

Table 6. Predicted and Validate Concentration of chromium at Different Depth.

<table>
<thead>
<tr>
<th>Depth [M]</th>
<th>Predictive Values</th>
<th>Experimental Values</th>
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<td>3.77E-06</td>
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Table 7. Concentration of Chromium at Different Depth.

<table>
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<th>Depth [M]</th>
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</thead>
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</tr>
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<tr>
<td>12</td>
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<td>3.83E-06</td>
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<td>45</td>
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</table>
The study has show the behaviour of the system through graphical representation expressing the behaviour of the chromium. The figures show the level of transport at different concentration. Figure one to four expresses the migration system under fluctuation phase, it implies that the deposition maintained high concentration at three metres to the lowest at thirty nine metres, these also express the rate of depositions in various strata structured in the study area, the concentration of chromium were observed to develop rapid heterogeneity on migration process within figure one and two, these increase are with respect to change in depth base on the transport system, it is influenced by the variation of the heterogeneity in porosity, such porous medium were observed to influences the migration rate of chromium concentration in the study area, while figure three maintained similar condition like one and two, fluctuation were observed with the optimum values recorded at sixty days. These developments increase the rate of concentration, because if the fluid velocity decreases its accumulation it definitely will reflect on the fluctuation increase in concentration; these conditions were experiences from figure one to four. While figure five and six maintained similar experiences, migration process observed fluctuation where the optimum values were recorded at twenty seven metres, but with different rate of concentration in the formation, figure seven and eight express it in physical process from high to low concentration, this can be attributed to slight deposition of inhibitors including rate of heterogeneity from degree of porosity between those strata. But slight increment of concentration were observed in figure seven and eight, the variation from porosity level and deposition of inhibitors were found to reflect on the concentration rate of chromium in these figures, their transport rate maintained fluctuation phase with increase in concentration from change in depth and time, their behaviour observed in this condition are base on the rate of inhibition and variations observed from increase rate of porosity in the formation, the transport of chromium were observed through these developed simulated values, the comparative analysis between predictive and experimental values generated best fits validating the developed model for chromium transport.

6. Conclusion

The study defined the behaviour of chromium in silt and fine sand formation, the study discusses the heterogeneity in various rate of concentration under the influences of porosity variation on its rate of deposition. The variation and inhibition observed in the transport process are reflected on the deposition of chromium in the study area. The migration rate of chromium has expresses its behaviour deposited on silt and fine sand formation, the developed model were to monitor the rate of concentration through its fluctuation phase expressed in graphical representation, the vacillation of the chromium in the strata shows the rate of influences from the stated parameters, the developed model were compared with experimental values, both parameters developed favourable fits, the behaviour of chromium has been expressed thoroughly through the developed model simulation values, the transport system in silt and fine sand formation has express the refection of slight immobile velocity generating slight fluctuation and accumulation in the study area.

References


