Application of superheated water as a soil remediation media: a review

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Abstract. Water is a unique solvent cause its highly hydrogen-bonded structure, and at room temperature it has a high boiling point for its mass, a high dielectric constant and high polarity. At the higher temperatures, its permittivity, viscosity and surface tension decreases, but diffusion rate increases. Superheated water is a general term to denote the region of the condensed phase between 100 °C and the critical point. Liquid water at elevated temperatures above its boiling point has been used for many years as an industrial solvent and cleaning agent in applications ranging from enhancing the extraction of oil shale, the extraction of sulphur from ore bodies in the Frasch process, to degreasing. In recent years, there has been an interest in using superheated water for soil remediation. In this paper, a review on this area of application has been performed.

1 Introduction

Unfortunately, one of the inseparable outcomes of rapid population growth are environmental pollutions. In many cases ignoring the polluted areas or regions at risk of pollution, will have irreparable effects on the environment and subsequently on the health of the human beings. In different areas in the country, especially the oil-rich ones, the local and sometimes wide-area pollutions due to the oil spill or carelessness in oil compounds transfer can be seen. Chemical industries, despite having so many benefits for providing welfare and comfort for the life of modern man can cause different diseases, including cancer, by entering the food cycle of humans; As in different studies, the carcinogenicity and mutagenetic effects of various hydrocarbon compounds have been proven. Soil, as the basis for producing needed foodstuffs for humans, is of great importance, and sometimes is ignored by this much. Entering various pollutants directly and indirectly (through the water and air) can cause some changes in this medium, and just like other elements of nature, the soil is always acting against these changes; But sometimes, the balance of the soil is disrupted, and we face the challenge of soil pollution. In these situations, human interventions in order to help the natural clearing processes and accelerating them seem necessary [1]. In other words, soils are considered as the refiners for nature. In addition to being a food supplier, soils also have purifying properties. These properties are originated from their physical properties (water diffusion through pores), chemical properties (adsorption and evaporation), and bio-based properties (decomposition and spoilage of organic matter). Contamination of soil with different chemical wastes is one of the most important environmental issues in various areas, which, due to the toxicity, carcinogenicity, and causing mutagenetic changes, there are vast concerns regarding their presence in nature [2]. These pollutants can be adsorbed on the surface of the soil or organic particles existing in it, which can enhance their concentration and subsequently cause their entrance to the surface waters along with the surface streams. On the other hand, these compounds may enter the groundwater and consequently lead to disorder in the food cycle of plants, humans, and animals, so they must be removed from the environment in some way [3]. Soil pollution factors include four major parts:

1. Agricultural factors
2. Oil contaminants
3. Industrial activities
4. Trash

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Regarding the soil pollution in Iran, according to the statistics, a few percent of industrial and domestic effluents are refined, and the major part of industrial and domestic effluents are discharged to the environment untreated. This manner can cause soil and groundwater pollutions, which increasing the nitrate content in well waters is a clear example for such consequences. One of the soil pollutions factors, which fortunately is not common in our country is acid rain. One of the actions of the department of environment to prevent acid rain is strong opposition to the establishment of a coke factory in the Savadkuh region. The atlas of soil pollution in Iran is being prepared, which provides accurate statistics on the amount of soil pollution in the country. Up now, the following regions are reported to be the most polluted regions in Iran: Isfahan province, south of Tehran, Asaluyeh in Bushehr, lands in the vicinity of the Mes-e Sarcheshmeh, Sistan and Baluchestan province, and Khuzestan province [4].

2 Modeling

The mass transfer mechanism of the extraction of soluble component is as follows: The soluble component attached to the solid network (by physical and chemical forces), have to be transferred to the solvent phase via liquidation or desorption. Then, the soluble component/solvent mixture diffuses to the solid surface and finally passes the static layer adjacent to the solid particles to reach the solvent phase. Since the contents of the solid particles vary with distance and time, the diffusion coefficients of the soluble component existing in the solid particles can be determined by observing the changes of concentration in the vicinity liquid with time. There are four mass transfer steps in general, which the diffusion of the soluble component in the solid particles toward the solvent is usually the determinative step of mass transfer [36].

Initially, there is a rapid extraction period with a constant intensity, which is stable in the mass transfer and controls the resistance intensity of the mass transfer film. Then, an unstable descending step with a very lower extraction intensity occurs. In this stage, the diffusion in the solid particles controls the intensity of the extraction process [36]. In industrial operating units such as extraction, a specific component in one phase diffuses to the contact point of two phases and then enters into the other phase. Transfer of one phase to the other phase continues until the two phases reach equilibrium point [37]. The results obtained by Fick’s second law were analyzed as a mathematical model. Since the modeling is performed in fluid phase, the following assumptions are considered:

1- Fluid moves with uz velocity.
2- Solid-phase is static.
3- There is no reaction.

In the fluid phase, movement is in z-direction, so the system is differentially modeled in this direction. According to the fluid stream, the diffusion term is ignored in this direction, and the convection term is considered as the determinative factor and described as follows:

\[ u_z \frac{c_{i,z}}{\partial z} \]

However, in radial direction, there is no movement, and lumped method is used for modeling, which only contains mass transfer flux:

\[ u_z \frac{c_{i,z}}{\partial z} \]

Now, the general relation of mass balance (input-output=accumulation) is shown below:

\[ k \alpha (c_i^* - c_{i,1}) + u_z \frac{c_{i,z}}{\partial z} + \frac{dc_{i,z}}{dt} = 0 \]

The initial and boundary conditions are as follows:

\[ c_{i,1}(0, t) = 0 \]
\[ c_{i,1}(z, 0) = 0 \]
\[ c_{i,1}(0, 0) = c_{i,1,0} \]

Changes are considered only in the z-direction, and concentration variations in other directions are neglected. The following image is showing the considered system for mass balance schematically.

By considering a cylindrical shape element with a base area equals to the tower base area and with a height equals to dz, the mass balance is written. At the element input, we have:

\[ N_{i,x} A \]

Where \( N_{i,x} \) and \( A \) are the i element flux in z-direction and tower base area, respectively. At the element output, we have:

\[ N_{i,x+dz} A \]

On the other hand, the mass transfer flux from the solid phase to gas phase is described as follows:

\[ (dz, A) k \alpha (c_i^* - c_i) \]

Where \( k \) is the mass transfer coefficient, \( \alpha \) is the specific surface area, and \( c_i \) and \( c_i^* \) are the i component concentrations in the liquid phase and solid phase surface, respectively. By substituting the mentioned equations on this base that the total mass transfer from the gas phase to the liquid phase is equal to output rate minus input rate plus accumulation rate, we have:

\[ \frac{\partial N_{i,x}}{\partial z} + \frac{\partial c_{i,z}}{\partial t} = k \alpha (c_i^* - c_i) \]

By dividing both sides of the above equation by dz,A, the following relationship is obtained:

\[ N_{i,x=0} A = N_{i,x} A + dz, A \frac{c_i}{\partial t} = (dz, A) k \alpha (c_i^* - c_i) \]

By considering the below equation:
N_i = u_i c_i + f_i

Furthermore, by defining J_i as follows:

J_i = -D_i \frac{\partial c_i}{\partial z}

In the above equations, u_i and D_i are the fluid velocity in the z-direction and the axial distribution coefficient. By substituting the above equations, we will have:

\frac{\partial (u_i c_i)}{\partial z} - \frac{\partial}{\partial z} \left( D_i \frac{\partial c_i}{\partial z} + \frac{\partial c_i}{\partial t} \right) = ka(c^*_{i} - c_i)

By assuming that the axial distribution coefficient does not depend on z-direction, we will have:

c_i \frac{\partial u_x}{\partial z} + u_x \frac{\partial c_i}{\partial z} - D_i \frac{\partial^2 c_i}{\partial z^2} + \frac{\partial c_i}{\partial t} = ka(c^*_{i} - c_i)

By considering the continuity equation and assuming that the fluid density is constant:

\frac{\partial u_x}{\partial x} + \frac{\partial u_y}{\partial y} + \frac{\partial u_z}{\partial z} = 0

Since the velocity in x and y directions are zero, the above equation turns into the following equation:

\frac{\partial u_x}{\partial z} = 0

Finally, we have:

\frac{\partial c_i}{\partial t} = D_i \frac{\partial^2 c_i}{\partial z^2} - u_x \frac{\partial c_i}{\partial z} + ka(c^*_{i} - c_i)

The above equation is the one to be solved. To solve such an equation, two boundary conditions and one initial condition are needed.

The boundary conditions are described as follows:

\frac{\partial c_i}{\partial z} = 0

The initial condition is determined as follows:

c_i(z, 0) = 0

In this study, the mass transfers of naphthalene, phenanthrene, fluoranthene, and pyrene from soil to the super-hot water were investigated. The information about molecular weight and solubility of these compounds are illustrated in the Table 1 [38].

c_i^* is the solubility of i component in groundwater. Therefore, given that the solubilities at 25 °C can be found in literature, the solubilities at 100, 150, 200, 250, and 300 °C should be calculated. To calculate the solubilities, the following equation is used [36]:

ln x_2(T) = \frac{T_0}{T} ln x_2(T_0) + 15 \left( \frac{T}{T_0} - 1 \right)^3

Where x_2(T) is the solubility at the temperature of T in the form of molar fraction and x_2(T_0) is the solubility at the specified temperature of T_0, and the temperature unit is Kelvin. To calculate the D_l, the following equation is used:

D_l = \frac{ud_p}{\varepsilon Pe}

Table 1. The information of the compounds used in this study.

| Compound       | Molecular weight | Solubility in water at 25 °C (mg/L) | Molecular volume (m³/kmol) |
|----------------|------------------|-------------------------------------|----------------------------|
| Naphthalene    | 2.128            | 32                                  | 407.0                      |
| Phenanthrene   | 2.178            | 3.1                                 | 554.0                      |
| Fluoranthene   | 3.202            | 2.0                                 | 6195.0                     |
| Pyrene         | 1.202            | 1.0                                 | 626.0                      |

Where d_p is the particle diameter, which is considered 2 mm in this study, the bed height is considered 10 mm, which is consistent with the empirical data, ε is the porosity coefficient (due to the presence of particles in soil), which is considered 0.4 in this work, and “u” is the entering velocity of the fluid. Four different flow rates, including 0.5, 1, 1.5, and 2 cm³/min, are considered in this work. Moreover, Pe is the dimensionless number of Peclet and can be calculated using the following equation [24]:

Pe = 1.634 Re^{0.265} S c^{-0.919}

According to the following equations:

Re = \frac{\rho u d_p}{\mu}

Sc = \frac{\mu}{\rho D_{ab}}

It is needed to know the viscosity and density at the specified temperatures. In order to determine these values, we use the equations below:

\rho = 858.03 + 1.2128 T - 0.0025 T^2

\mu = \exp(-10.2 + \frac{2990970}{T^2})

The units of temperature, density, and viscosity in the above equations, are Kelvin, kg/m³, and Pa.S, respectively.

To calculate the diffusion coefficient (D_{ab}), the following equation can be used:

D_{ab} = \left(117.3 \times 10^{-18}\right) (\phi M_p)^{0.5} T

Where M_p is the molecular weight of solvent (water in this case), \phi is the correlation coefficient of solvent, which is equal to 2.26, and V_a is the molar volume of the soluble, which can be obtained from \nu_e and can be seen in the Table 1.

\nu_e = 0.285 (\nu_e)^{1.048}

To calculate the mass transfer coefficient (k), the dimensionless number of Sherwood (Sh) is used:

k = \frac{Sh . D_{ab}}{d_p}

Where the Sherwood number can be obtained from [24]:

Sh = 0.38 Re^{0.83} Sc^{1/3}
3 Results and discussion

3.1 Time effect

The extraction time in the method of extraction with superhot water is short. To extract compounds from plant samples or investigate the environmental samples, in 5 min to 2 hr in a dynamic state and 5-10 min in a static state, suitable efficiency can be achieved [3]. By increasing the extraction time, the efficiency will increase up to a constant value. The extraction time depends on the kinetics of the process and the time needed to reach the equilibrium. It is expected that by increasing the extraction time, the efficiency increases. The effect of time on the extraction of hydrocarbons or multiple-ring aromatic is shown in Figures (1,2).

Fig. 1. Changes of fluoranthene concentration along the tower at different times at 373 K.

As it is obvious, by increasing the extraction time, the concentrations of hydrocarbons in the liquid phase are increased. In fact, from a specific time to further, there is no change in the curve by increasing the time, which means it reached equilibrium. Also, it can be concluded that by increasing the time, the change rate of concentration increases, which can be ascribed to the contact time of two phases. In other words, the sufficient time needed for mass transfer is provided by further increasing the extraction time.

Fig. 2. Changes of pyrene concentration along the tower at different times at 373 K.

3.2 Flow rate effect

The intensity of extraction enhances by increasing the flow rate. It is expected that a more flow rate can mitigate the resistance of the fluid phase. Thus, the amount of mass transfer from the solid phase to the fluid phase increases. In the following figures (Figures (3-6)), the effect of changes in flow rate in the range of 0.5-2 cm³/min is shown for the four mentioned compounds.

Fig. 3. The effect of flow rate on the concentration of phenanthrene.

Fig. 4. The effect of flow rate on the concentration of fluoranthene.

Fig. 5. The effect of flow rate on the concentration of naphthalene.

As can be seen in Figures, the variation of flow rate is not determinative on the amount of concentration. Also, for various compounds, there is no significant change in concentration, which can be concluded that the main resistance against mass transfer is the internal resistance, no
the external one. In other words, the diffusion in the solid phase and reaching to surface, feel the most resistance. Overall, the resistance of the liquid phase is negligible.

Fig. 6. The effect of flow rate on the concentration of pyrene.

3.3 Temperature effect

The most important parameter that affects the super-hot water system is temperature because by changing the temperature of the solvent (water), its properties will change [11, 39, 40]. By increasing the temperature, the surface tension and viscosity decrease, whereas the diffusion of the soluble compounds in the water enhances. In other words, more sufficient mass transfer will be provided by increasing the temperature, which can improve the extraction efficiency. This phenomenon is even more important in weak-polar or non-polar compounds because their solubility in water is negligible in low temperatures. Temperature can affect the solubility in water, axial distribution coefficient, and diffusion coefficient. So, significant changes in concentration are expected by varying the temperature. At the 25°C temperature, as can be seen in the previous section, water has a weak capability to dissolve these kinds of hydrocarbons. As the temperature increases, the dielectric constant decreases, and consequently, the solubility increases. Therefore, it is expected that by increasing the temperature, the solubility of this group of hydrocarbons in the liquid phase enhances.

Fig. 7. The effect of temperature on the concentration of phenanthrene.

As shown in Figures, by increasing the temperature ranging from 200 to 250, the concentration variations were significantly affected. This phenomenon can be assigned to the effect of these compound’s solubility, which by increasing the temperature, the solubility enhances significantly, and this is obvious in Figures.

Fig. 8. The effect of temperature on the concentration of fluoranthene.

Fig. 9. The effect of temperature on the concentration of pyrene.

Fig. 10. The effect of temperature on the concentration of naphthalene.
The following figure provides a comparison between the concentration changes of the four compounds at 2 cm³/min flow rate and 150 °C temperature. As can be seen in Figure, the concentration changes of naphthalene are more than others, and phenanthrene, fluoranthene, and pyrene come next, respectively. This order exactly originates from their solubility which is in the order of naphthalene, phenanthrene, fluoranthene, and pyrene, from higher to lower. On the other hand, the molecular weight of naphthalene is lower than others, which influences on its diffusivity in the liquid phase.

3.3.1 Review of experimental studies

This review is done for the compounds of naphthalene and phenanthrene [41].

The concentrations of remaining naphthalene were 29, 1.8, 1.9, 0, and 0 ppm, at 100, 150, 200, 250, and 300, respectively. This means that the concentration of remaining naphthalene at 100 degrees, is slightly more than its concentration in the initial soil, and at 150 degrees, is significantly decreased to a value of 2 ppm. This evidences that the extracted naphthalene does not separate from soil in 100 °C. But, over 150 °C, it will be easily separated from the soil. There are some other reasons that why naphthalene does not separate from the soil at 100 °C. One of them is that the physical properties of soil up to 100 °C temperature and 100 atm pressure will change, and this variation can cause a change in the separation fraction of methanol from the soil and subsequently will affect the separation of remaining naphthalene from the soil. As another reason, the lack of homogeneity in the concentration of initial soil, which is heated up to 100n °C, can be mentioned. The concentration of remaining naphthalene was also decreased from 30 ppm at 100 °C, to 0.74 ppm at 300 °C. As can be seen in Figure 2, the separation of phenanthrene from the soil needs more heat than the value needed for naphthalene, which originates from the properties of phenanthrene that, in comparison with naphthalene, has a lower solubility in water and higher molecular weight and melting point.

To clear a naphthalene-contaminated soil, 150 °C is sufficient. To clear a soil polluted with PAHs such as phenanthrene, anthracene, etc.

4 Conclusions

No need for organic solvent or negligible assumption of organic solvent is the most prominent characteristic of extraction with super-hot water. Accessibility, low-cost, and non-toxicity of water as the main solvent, and no need for drying the wet samples are other characteristics of this method. In many cases, this method in comparison with other methods, is more clear, cheaper, and faster. Extraction with super-hot water has showed lots of benefits in term of no need for organic solvents in environmental extractions. The equipment needed for this method is relatively simple and there is no need for applying high pressures, as the extraction with supercritical fluid. In this study, a simple mathematical model was presented for the fluid phase, which evaluated the extraction efficiency in terms of temperature, time, and flow rate. Also, the amount of concentration changes for naphthalene, phenanthrene, fluoranthene, and pyrene were measured with the parameters of temperature, flow rate, and time. Among these parameters, temperature was the most determinative one, and the amount of extraction is increased by increasing the temperature. It can be mentioned that almost 95% naphthalene was extracted at 100 bar. Because naphthalene
was resistant in soil at 100 °C and does not separate from the soil, but at 150 °C and higher temperatures, due to its physical properties, which is changed in this condition, it can be easily separated from soil. As in this study, significant changes in concentration were observed in the temperatures of 200 and 250 °C, which is ascribed to the effect of solubility of these compounds that increases by increasing the temperature. Also, the variation of flow rate has a negligible effect on the concentration for all compounds, which means that the internal resistance is determinative, comparing with the external one. In other words, the diffusion in the solid phase and reaching to the surface feel the most resistance. Overall, the resistance of the liquid phase is negligible. By increasing the extraction time, the rate of concentration changes increases, which can be assigned to the contact time between two phases. By a meaning, sufficient time for mass transfer can be provided by increasing the time. Short extraction time, high-quality extract, low cost of indicators needed for extraction, and compatibility with the environment are other prominent characteristics of this method.

5 Limits and suggestions

According to this fact that the solvent used in the extraction with super-hot water is water, the more assumption of water comparing with other methods can be mentioned as one of the disadvantages of this method, which is suggested that to overcome this issue in future studies, this should be included in the technical and economical calculations.

Also, as one of the limits of this method, high temperature and pressure can be mentioned, which can be a serious drawback regarding utilizing this method in more wide applications, so it is needed to be studied more.

Corrosion is also another important factor that should be considered. For example, stainless steel, as one of the most applicable material that is usually used in the equipment of super-hot water is corroded. So, for dealing with super-hot water, utilizing special materials is needed, which are more expensive than common materials in the industry. According to the discussed contents, it is hoped that the existing problems to be solved or mitigate in future studies, to facilitate the more application of this technology in industry.

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