Computational simulation and modelling of uranium extraction using tributylphosphate through membrane extractor

Rahmad Syah1, Dadan Ramadan1, Marischa Elveny2*, Yan Cao3*, Afrasyab Khan4, Hamid Abdi5 & Mahdi Ghadiri6

Non-disperse solvent extraction is an effective technique for the extraction of metal ions from aqueous solution. In this study, uranium extraction using n-dodecane solvent containing tributylphosphate extractant in a membrane contactor was investigated. A 2D mathematical model was developed for the fluid flow and mass transfer in the hollow fibre membrane extractor. The equations of the created model were solved using the finite element method. The uranium concentration distribution in the extractor at different extractant concentrations as well as feed acidity was studied. The results showed that there is reasonable good agreement between experimental uranium extraction and modelling outputs at different extractant concentrations. Increasing extractant concentration from 5 to 30% led to the enhancement of uranium extraction from 2.60 to 34.13%. Also, there was an increase in the uranium extraction with increasing feed acidity in the range of 1–3 M. Furthermore, based on the radial uranium concentration distribution, it was found that the main mass transfer resistance in the system was microporous membrane section. Finally, it was obtained that the uranium extraction efficiency could be improved significantly by increasing porosity-to-tortuosity ratio. It was concluded that the membrane specification plays the most important role as the dominant mass transfer resistant was in the membrane subdomain.

Uranium is considered as the main source for production of nuclear power and radiopharmacy1. It seems that the uranium consumption in the world will be increased to 82,195 tons by 20252. There are various type of uranium resources including uranium-containing complex ores3, copper and phosphates rocks4, low-grade uranium ore and tailings containing uranium1.

After leaching or bioleaching, there are different downstream processes for uranium recovery and purification including direct precipitation, ion-exchange (IX), as well as solvent extraction (SX). Among these methods, the solvent extraction is common due to the high costs of other methods5. Uranium microfluidic solvent extraction was done using the organic phase containing 30% (v/v) tributyl phosphate as extractant in dodecane. Using two-stage microfluidic extraction approximately 90% uranium extraction was obtained at contact time of less than 3 s6. Di-(2-ethylhexyl) phosphoric acid (D2EHPA) has been used as effective extractant for the uranium extraction. The amount of 1 M D2EHPA extracted about 94% of uranium using two-stage extraction7. There was 99.4% uranium extraction when mixture of Primene JM-T as well as Alamine 336 was used as extractants8. Despite of many benefits of the conventional solvent extraction, it has a number of shortcomings including coalescence and density difference between aqueous and organic solutions, emulsification issue, require for dispersion, flooding, and etc. In order to overcome these issues, the non-dispersive solvent extraction (NDSX) process has been introduced as an alternative approach. In NDSX, a porous membrane is placed between two phases and the aqueous–organic contact with each other is happened on the surface of membrane interface. So,
the separation of phases is not needed in this process and the flooding and loading issues do not observe due to independent flow of aqueous and organic phases.\textsuperscript{9–14}

Uranium was extracted form nitric acid feed solution using different amides extractants in a membrane contactor, then, the stripping process was performed in a separate membrane contactor to transfer uranium from organic solution to HNO\textsubscript{3} or Na\textsubscript{2}CO\textsubscript{3} solution.\textsuperscript{15} Ansari et al.\textsuperscript{16} used non-dispersive solvent extraction for the uranium recovery using different extractants include D\textsubscript{2}EHPA, N,N-diethyl decanamide (DHDA), N,N-diethyl hexanamide (DHHA), TBP and N,N-diethyl octanamide (DHOA). The results showed that monoamide extractants have good capability for the extraction of uranium. There have been a number of studies on the experimental studies of uranium NDSX extraction.\textsuperscript{15,16} However, it is incredibly difficult to find works which the simulation and modelling of uranium extraction were investigated. Swain et al.\textsuperscript{17} used a computational fluid dynamics (CFD) mathematical model to study U extraction from HNO\textsubscript{3} medium by tri-n-butyl phosphate (TBP) as extractant in n-dodecane as dilute in a hollow-fibre membrane contactor (HFMC). Effect of different operating parameters on the extraction percentage were evaluated. They used Navier–Stokes equations to find velocity profile in the shell side.

The Swain et al.\textsuperscript{17} work was extended in the current study in order to examine in more details effect of various critical parameters of diffusive and convective uranium flux in the membrane contactor. Moreover, the change in diffusion coefficient and physical properties like viscosity as well as density with change in extractant concentration was considered. Also, in this study Happel’s free surface model\textsuperscript{18} was used to determine velocity profile in the shell compartment of contactor.

Numerical investigation

Uranium extraction is took place by two various mechanisms including diffusion as well as convection along the membrane contactor. The continuity equation is considered as the most important equation that explains uranium transfer phenomena in three subdomains. Three subdomains were shown in Fig. 1. Furthermore, the reaction mechanisms can be written as follow:\textsuperscript{17,19}

\[
UO_2^{2+} + 2NO_3^{-} + 2TBP \leftrightarrow UO_2(NO_3)_2 \cdot 2TBP_{\text{org}}
\]  

(1)

For reaction (1), the U extraction equilibrium constant, \( K_{\text{ex}} \), is determined as follows:

\[
K_{\text{ex}} = \frac{UO_2(NO_3)_2 \cdot 2TBP_{\text{org}}}{UO_2^{2+} + 2NO_3^{-} + 2TBP}
\]

(2)

The continuity equation can be written as follows at steady state conditions. There is no reaction in the tube and shell sides (Eq. 3) and membrane domain (Eq. 4):\textsuperscript{20,21}

\[
\text{div}(C_{\text{Uranium}} V_z + J_{\text{Uranium}}) = 0
\]

(3)

\[
\text{div}(J_{\text{Uranium}}) = 0
\]

(4)
where $C_{\text{Uranium}}$, $J_{\text{Uranium}}$, and $V_z$ refer the concentration of uranium (mol m$^{-3}$), diffusion flux and the velocity symbol (m/s) respectively. For the diluted solutions, the Fick's law is usually used for the calculation of solute transfer. Therefore, the diffusion flux of uranium can be written as follows and it can be used in equations of (3) and (4)\(^{(20)}\):

$$J_{\text{Uranium}} = -D \nabla C_{\text{Uranium}}$$  \hspace{1cm} (5)

It could be possible to ignore the radial convective flux because the solution velocity is tiny in $r$ direction. Therefore, the mass transfer equations for the tube and shell subdomains are written as\(^{(20)}\):

$$D_{\text{Uranium,tors}} \left[ \frac{\partial^2 C_{\text{Uranium,tors}}}{\partial r^2} + \frac{1}{r} \frac{\partial C_{\text{Uranium,tors}}}{\partial r} + \frac{\partial^2 C_{\text{Uranium,tors}}}{\partial z^2} \right] = V_{z,tors} \frac{\partial C_{\text{Uranium,tors}}}{\partial z}$$  \hspace{1cm} (6)

In membrane subdomain, there is only diffusive flux for uranium transfer, so, the convective flux term needs to be removed from the equation\(^{(20)}\):

$$D_{\text{Uranium,m}} \left[ \frac{\partial^2 C_{\text{Uranium,m}}}{\partial r^2} + \frac{1}{r} \frac{\partial C_{\text{Uranium,m}}}{\partial r} + \frac{\partial^2 C_{\text{Uranium,m}}}{\partial z^2} \right] = 0$$  \hspace{1cm} (7)

Diffusion coefficient in the microporous membrane pores was calculated using Eq. (8)\(^{(22)}\):

$$D_{\text{Uranium,m}} = D_{\text{Uranium, organic phase}} \frac{r}{\tau}$$  \hspace{1cm} (8)

Membrane subdomain boundary condition may be expressed as follows:

$$r = R_1, C_{\text{mem}} = C_{\text{tube}} \times m(\text{concentration})$$  \hspace{1cm} (9)

$$r = R_2, C_{\text{mem}} = C_{\text{shell}}(\text{concentration})$$  \hspace{1cm} (10)

The aqueous phase containing uranium velocity in the tube in the axial direction is determined using parabolic laminar flow as\(^{(16,23)}\):

$$V_z = 2 u_t \left[ 1 - \left( \frac{r}{R_1} \right)^2 \right]$$  \hspace{1cm} (11)

The symbol $u$ means velocity of aqueous phase (m/s), and $R_1$ is the fibre inside radius (m). The boundary conditions for uranium mass transfer can be given as follows\(^{(16,24,25)}\):

$$Z = 0, \text{ for lumen subdomain } \rightarrow C = C_{\text{inlet}}(\text{initial concentration of uranium})$$  \hspace{1cm} (12)

$$Z = L, \text{ for lumen subdomain } \rightarrow n(-D \nabla C_{\text{Tyramine}}) = 0(\text{Convective flux})$$  \hspace{1cm} (13)

$$r = 0, n \cdot (-D \nabla C_{\text{Tyramine}} + C_{\text{Tyramine}} V_z) = 0(\text{axialsymmetric})$$  \hspace{1cm} (14)

$$r = R_1, C_{\text{tube}} = C_{\text{mem}}/m(\text{concentration})$$  \hspace{1cm} (15)

The velocity of organic solution containing extractant in the shell in the axial direction is calculated using Happel’s free surface model\(^{(20)}\):

$$V_z = 2 u_t \left[ 1 - \left( \frac{R_2}{R_3} \right)^2 \right] \times \left[ \frac{\left( \frac{R_2}{R_3} \right)^2 - \left( \frac{R_3}{R_2} \right)^2 + 2 \ln \left( \frac{R_3}{R_2} \right)}{3 + \left( \frac{R_2}{R_3} \right)^4 - 4 \left( \frac{R_3}{R_2} \right)^2 + 4 \ln \left( \frac{R_3}{R_2} \right)} \right]$$  \hspace{1cm} (16)

Also, the boundary conditions for uranium mass transport can be given as follow\(^{(21)}\):

$$Z = 0, n(-D \nabla C_{\text{Tyramine}}) = 0(\text{Convective flux})$$  \hspace{1cm} (17)

$$Z = L, \text{ shell side } \rightarrow C = C_{\text{inlet}}(\text{inlet concentration})$$  \hspace{1cm} (18)

$$r = R_2, C_{\text{shell}} = C_{\text{mem}}(\text{concentration})$$  \hspace{1cm} (19)

$$r = R_3, n \cdot (-D \nabla C_{\text{Tyramine}} + C_{\text{Tyramine}} V_z) = 0(\text{Insulation})$$  \hspace{1cm} (20)

Table 1 lists the extractor’s parameters, fluid characteristics, and operational conditions.

**Numerical solution.** The obtained equations for the numerical investigation of uranium transport through membrane were solved via COMSOL Multiphysics software at steady state condition. The equations, geometry, boundary conditions were defined in the software. Then, it was solved after meshing the geometry. The meshing of membrane extractor was shown in Fig. 2. The mesh element size in the membrane subdomain and around
Table 1. The extractor’s parameters, as well as the characteristics of the fluids and the operating conditions\textsuperscript{17,26}.

| Parameters                      | Unit  | Values          |
|---------------------------------|-------|-----------------|
| Uranium initial concentration   | g/L   | 21.5            |
| Feed flow rate                  | Lph   | 10–60           |
| Organic phase flow rate         | Lph   | 10–50           |
| TBP concentration (\%)          |       | 5–40            |
| Feed acidity                    | M     | 1–4             |
| Temperature                     | K     | 296             |
| Fibre inside diameter           | m     | 2.4e⁻⁴          |
| Shell inside diameter           | m     | 63.6            |
| Effective length of tube (L)    | m     | 0.15            |
| Fibre thickness                 | m     | 3e⁻⁵            |
| Porosity (ε)                    | vol%  | 0.3             |
| Tortuosity (τ)                  |       | 3.75            |
| Number of fibres                | –     | 10,000          |
| Operation mode                  | –     | Counter-current, once through |

Figure 2. The generated mesh in the membrane extractor.
its surface was built smaller where the complex between uranium ion and TBP is formed. It was performed to increase the accuracy of the developed model. The mass transport equations were solved using the UMFPACK numerical solver as a linear solution. To solve the created model and obtain the results, a machine with RAM of 8.00 GB, Intel Core™ i7 CPU, and 64-bit operating system was applied.

Mesh independence test is required to perform in any CFD models to make sure that the solution is also independent of the mesh resolution. Influence of element numbers on the extraction of uranium is shown in Fig. 3. It was observed that increasing element numbers from 322 to 15,160 led to the decreasing of uranium extraction from 35.91 to 34.11%. It was not observed meaningful change in uranium extraction after 7042 number of elements. Also, unnecessary increasing of mesh number can cruise increasing of the cost and time of simulation. Therefore, elements number of 7042 was selected as the optimum value.

Results and discussion

Model validation. The amount of uranium extraction at different TBP concentration was shown in Table 2. The experimental results provided in Table 2 are those reported Swain et al. Increasing in extractant concentration results in the enhancement of uranium extraction from aqueous solution in to organic phase. This increase is because of uranium distribution coefficient increasing with increasing TBP concentration. Furthermore, as it can be observed there is reasonably great agreement between experimental values of uranium extraction and modelling outputs. Only, small difference was observed at the highest TBP concentration but generally it was seen that the developed model is able to predict the uranium extraction accurately.

Uranium concentration distribution. Uranium concentration distribution and its total flux including diffusive and convective fluxes are presented in Fig. 4. The uranium concentration decreases along membrane contactor at the tube side. It is because a complex is formed between uranium ion and TBP on the membrane surface, then, it penetrates through pores of microporous membrane. At the entrance, uranium diffusion flux is high where the concentration gradient between two phases is larger. The amount flux is higher at the centre of tube and shell sides where the velocity is maximum while the total flux is low around membrane surface where the velocity is minimum and the main flux mechanism is diffusive flux. Also, it should be pointed out that the diffusive flux is horizontal and in favour of uranium extraction. It was seen in the membrane subdomain where the only flux is diffusive flux.

TBP concentration effect on the uranium axial and radial concentration distribution. Figures 5 and 6 show the dimensionless uranium concentration distribution in axial direction at tube side and in

| TBP concentration (%) | Exp. U extraction | Model U extraction |
|-----------------------|------------------|--------------------|
| 5                     | 2.5              | 2.59               |
| 10                    | 5                | 4.44               |
| 20                    | 16               | 16.83              |
| 30                    | 29               | 34.13              |

Table 2. Comparison of experimental data and modelling outputs in terms of uranium extraction.
radial direction in the contactor at different TBP concentration. According to Fig. 5, the uranium concentration decreases along membrane at the tube side due to extraction of uranium into the organic solution in the shell subdomain of contactor. Furthermore, increasing TBP concentration can result in more decreasing in uranium concentration at the outlet of tube side. Based on the Fig. 6, it can be seen from that the values in the tube and shell sides are roughly the same (between 0.8 and 1 at tube side at different TBP concentration) and (between around 0.005 and 0.02 at shell side at different TBP concentration). However, there is significant drop in the membrane side especially for higher TBP concentration. It could be concluded that the main resistance for the uranium mass transfer is in the membrane contactor.

**Feed acidity effect.** Solution pH is one of the important parameters in non-dispersive solvent extraction as it can have considerable effect on the formation of complex between target ions in aqueous phase and extraction molecule in the organic phase and subsequently on the target ion partition coefficient. Uranium dimensionless concentration distribution as a function of axial distance was provided in Fig. 7. The modelling outputs clearly show that the concentrations of uranium go down along membrane contactor. Also, it was observed that the differences of higher feed acidity is much larger than that of lower acidity. This reveals that the concentration gradient of uranium is much larger at higher feed acidity because of larger partition coefficient at higher feed acidity. Also, uranium dimensionless concentration distribution as a function of radial distance in the middle of membrane contactor at different feed acidity was shown in Fig. 8. It can be clearly observed that the uranium concentration distribution at tube and shell sides are almost horizontal straight line. The concentration gradient is higher for the feed with higher acidity at the membrane subdomain of contactor. The maximum concentration on the membrane surface at tube side was obtained 3.55 mol m\(^{-3}\), 5.31 mol m\(^{-3}\), and 6.05 mol m\(^{-3}\) for the feed acidity of 1, 2, and 3 molar. The uranium concentration was decreased to 0.018 mol m\(^{-3}\), 0.016 mol m\(^{-3}\), and 0.11 mol m\(^{-3}\) on the membrane surface at the shell side of contactor for the feed acidity of 3, 2, and 1 molar.

![Figure 4. Uranium concentration distribution and its total flux.](image-url)
Based on this large drop in uranium concentration, it could be concluded that the main resistance for uranium mass transfer is at the microporous membrane. Influence of feed acidity on the uranium extraction is shown in Fig. 8. According to Fig. 9, the uranium extraction was increased from 9.37 to 33.17% with the enhancement of feed acidity from 1 to 3 M. In fact, decreasing in outlet uranium concentration (Fig. 7) means that the more uranium was transferred into shell side and the uranium extraction was increased.

**Effect of porosity-to-tortuosity ratio on uranium extraction.** Based on “TBP concentration effect on the uranium axial and radial concentration distribution” and “Feed acidity effect”, it was found that the main mass transfer residence for uranium transport through membrane belongs to the membrane subdomain. Also, it is observed from Eq. (6) that the membrane porosity and tortuosity has a considerable influence on the mass transport of uranium. A higher porosity and lower tortuosity will lead to decreasing of membrane mass transfer resistance\(^{22}\). In the current study, the influence of membrane porosity-to-tortuosity on uranium extraction was studied and provided in Fig. 10 using two different equations (structure factor)\(^{22,27}\).

\[
\tau = \frac{1}{\varepsilon} \tag{21}
\]

\[
\tau = \frac{(2 - \varepsilon)^2}{\varepsilon} \tag{22}
\]

Based on the Eqs. (19) and (20), there is also direct relationship between the membrane porosity and its tortuosity. The enhancement of the membrane porosity will result in the decrease in tortuosity. Increasing porosity from 0.15 to 0.8 was increased the uranium extraction from 11.4 to 92.73% and 3.51–86.02% according to Eqs. (21) and (22), respectively. From the model outputs, it was found that the membrane mass transfer resistance increases with the decreasing of porosity. As in the uranium extraction system the main mass transfer resistance is in the membrane subdomain, it can be reduce by increasing the membrane porosity. But, in terms of membrane manufacturing, Because of fabrication and mechanical strength concerns, there are certain limitations on membrane porosity increasing.
Conclusion

A mathematical model was proposed with the use of finite element method to investigate the transport of uranium ions through the microporous membranes. Non-disperse solvent extraction system was divided into three compartments including tube, membrane, and shell. The aqueous solution containing uranium flows at the tube side while the organic phase containing TBP enters the shell side counter-currently. The developed model solved the mass transfer equations for the prediction of uranium concentration distribution and its extraction in the contactor. According to the results, increasing extractant concentration and feed acidity had positive impact on the uranium extraction. Moreover, it was found that the main mass transfer resistance was membrane section and it was observed a significant drop in the uranium concentration in this section. At the end, uranium extraction as a function of porosity-tortuosity ratio was investigated and it was seen that there is considerable enhancement in the uranium extraction with increasing this ratio. Increase in this ratio can result in increase in uranium diffusion coefficient in the membrane pores.
Figure 7. Dimensionless uranium concentration distribution along membrane contactor at different feed acidity.
Figure 8. Dimensionless uranium concentration distribution in radial direction at different feed acidity.
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Figure 9. Change in uranium extraction as a function of feed acidity.

Figure 10. Uranium extraction as a function of porosity-to-tortuosity ratio.
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Author contributions

R.S: Validation, Modelling, Revising manuscript, Applying reviewers commentsD.R: Validation, Revising manuscript, Applying reviewers commentsM.E.: Resources, Software, Reviewing and Editing, Data curation, Revising manuscriptY.C.: Software, Revising manuscript, Applying reviewers commentsA.K: Conceptualization, Methodology, Formal analysis, Modelling, Writing Original draft preparationH.A.: Conceptualization, ResourcesM.G.: Conceptualization, Resources, Reviewing and Editing, Supervision, Project administration.

Competing interests

The authors declare no competing interests.

Additional information

Correspondence and requests for materials should be addressed to M.E. or Y.C.

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