Abstract. The dissolution kinetics of potassium sulfate dissolved in water under pneumatic mixing has been investigated. The mass transfer coefficients were determined by the experiments, as well as on the basis of the local isotropic turbulence theory. The ratio between theoretical and experimental data was found to be satisfying that allows to determine the theoretical mass transfer coefficient to calculate the mentioned process.

Keywords: dissolution, kinetics, mass transfer coefficient, intensification.

1. Introduction

Dissolution processes are used in many branches of national economy, therefore they are of great importance [1]. Chemical, food, hydrometallurgical, pharmaceutical, galvanic and other industries are interested in studying such processes [2-5]. The data of dissolution process statics and kinetics are the basis for its calculations and designing. In accordance with mentioned data the equilibrium concentrations, degree of equilibrium and time of technological dissolution process are determined. To determine the dissolution time it is necessary to know the kinetic coefficients, namely mass transfer coefficient, which is determined under the laboratory conditions. Commonly the data about mass transfer coefficients are generalized and provided as dimensionless complexes of similarity criteria. The basis of such generalizations is the similarity theory and kinetic results of dissolution in the criteria forms are represented as Eq. (1):

\[ Sh = f(Re, Ar, Sc...) \] (1)

where \( Sh = \frac{\beta \cdot 1}{D} \) – Sherwood number; \( Re = \frac{\nu \cdot 1}{\nu} \) – Reynolds number; \( Ar = \frac{g \cdot 1^2 \cdot \Delta \rho}{\nu^2 \rho} \) – Archimedes number; \( Sc = \frac{\nu}{D} \) – Schmidt number; \( \beta \) – mass transfer coefficient, m/s; typical linear size, m; \( D \) – diffusion coefficient, m^2/s; \( \nu \) – velocity of liquid flowing around the solid body, m/s; \( \nu \) – kinematic viscosity coefficient, m^2/s; \( \rho \) – liquid density, kg/m^3; \( \Delta \rho \) – difference between solid body and liquid densities, kg/m^3.

Archimedes number is used in such a case when it is impossible to determine the value \( \nu \).

Opening of the function in Eq. (1) is given by many investigators, e.g. by R. Petrus et al. [6].

Another method is a theoretical analysis based on the presence of developed turbulence in liquid which is characterized by the turbulence scales from large to small ones [7]. Small-scale turbulence is characterized by homogeneity and isotropy. The mass transfer coefficient \( \beta \) is determined based on these properties for the dissolution processes under the mechanical stirring in the solid body-liquid system [8]:

\[ \beta = 0.267(e_0 \nu)^{0.25} Sc^{-0.75} \] (2)

where \( e_0 \) – the average value of the energy refered to the mass unit of the system, W/m^2.

Under the mechanical stirring the energy is determined on the basis of that energy introduced into the system by the mechanical stirrer.

During our investigations the stirring is carried out by a gas phase which considerably changes the system physical parameters due to the formation of liquid-gas emulsion. Momentary isolation of solid particles by gas bubbles may have a positive effect due to the non-stationarity of the dissolution process on the separate particles. Using gas phase has advantages in the case of corrosive medium, dissolution in large volumes and uniform stirring of the whole system.

The aim of this work is to investigate the kinetic regularities of monofractional solid phase dissolution in liquid stirred by gas; experimental determination of mass transfer coefficients taking into account the change of motive force and particles size; comparison of the determined coefficients with \( \beta \) values determined on the basis of local isotropic turbulence.
2. Experimental

2.1. Theoretical Part

The material balance of dissolution determines the interaction between the solid body and salt masses in the solution and has the form:

\[ M_0 - M = V \cdot c \]  

(3)

where \( M_0 \) – the salt initial mass, kg; \( M \) – the alt mass at dissolution time \( t \), kg; \( V \) – liquid volume, \( m^3 \); \( c \) – the salt concentration at time \( t \), kg/m\(^3\).

Eq. (3) corresponds to the condition when the salt initial concentration is equal to zero in liquid. Introducing the relative values \( \eta = \frac{M}{M_0} \) and \( c_{\text{max}} = \frac{M_0}{V} \) we obtain Eq. (4), the equation of a working line, which is a straight dependence

\[ \eta = 1 - \frac{1}{c_{\text{max}}} \cdot c \]  

(4)

It determines a degree of deviation from equilibrium value in liquid \( c_e \).

The process kinetics is determined by the following equation [1]:

\[ -\frac{dM}{dt} = \beta \cdot A \cdot (c_e - c) \]  

(5)

where \( A \) – dissolution surface, \( m^2 \) and \( t \) – time, s.

The values \( M \), \( A \) and \( t \) are variable in Eq. (5). Using Eq. (4) let us express the concentration by means of the system linear sizes and take into consideration that for the spherical particles

\[ \eta = \frac{M}{M_0} = \frac{d^3}{d_0^3} = \varphi^3 \]  

(6)

running values of the solid body mass are equal to

\[ M = N \cdot \rho_s \cdot \frac{\pi}{6} \cdot d^3 \]  

(7)

and for the surface

\[ A = N \cdot \pi \cdot d^2 \]  

(8)

where \( N \) – particles number; \( \rho_s \) – the solid body density, kg/m\(^3\).

Substituting (7) and (8) in Eq. (5) and taking into account the value from (6), after transformations we obtain Eq. (9):

\[ -\frac{d(d)}{dt} = 2 \cdot \frac{\beta \cdot c_{\text{max}}}{\rho_s} \left[ \left( \frac{c}{c_{\text{max}}} - 1 \right) + \varphi^3 \right] \]  

(9)

To simplify the integration of Eq. (9) we introduce dimensionless linear size \( \varphi \); the value \( \left( \frac{c}{c_{\text{max}}} - 1 \right) \) we denote as \( a^3 \). The Eq. (9) has the form:

\[ -\frac{d \left( \frac{\varphi}{a} \right)}{dt} = 2 \cdot \frac{\beta \cdot c_{\text{max}} \cdot a^2}{\rho_s \cdot d_0} \left[ 1 + \left( \frac{\varphi}{a} \right)^3 \right] \]  

(10)

Integration of Eq. (10) within the limits: time \( t \) from 0 to \( t \) and \( \frac{\varphi}{a} \) from \( \frac{1}{a} \) to \( \frac{\varphi}{a} \) allows to determine the dissolution kinetics of solid particles in the integral form [9]:

\[ \phi \left( \frac{1}{a} \right) \phi \left( \frac{\varphi}{a} \right) = \frac{c_{\text{max}} \cdot a^2 \cdot 2 \cdot \beta t}{d_0} \]  

(11)

where functions \( \phi \left( \frac{1}{a} \right) \) and \( \phi \left( \frac{\varphi}{a} \right) \) and \( \phi \left( \frac{1}{a} \right) ma \phi \left( \frac{\varphi}{a} \right) \) are determined by the following dependence [9]:

\[ \phi(x) = \frac{1}{6} \ln \left( \frac{1 + x^2}{1 - x + x^3} \right) + \frac{1}{\sqrt{3}} \arctg \frac{x\sqrt{3}}{2 - x} \]  

(12)

Let us substitute \( \frac{1}{a} \) or \( \frac{\varphi}{a} \) in Eq. (12) instead of \( x \).

After we plotted \( \phi \left( \frac{1}{a} \right) \phi \left( \frac{\varphi}{a} \right) = f(t) \) we determined the mass transfer coefficient \( \beta \) by means of a slope angle.

The aim of this work is to compare the calculated value with the mass transfer coefficient \( \beta \) obtained on the basis of local isotropic turbulence (Eq. (2)). Gas phase provides more uniform stirring of liquid compared with the mechanical stirrer due to the uniform distribution of gas along the apparatus cross-section.

2.2. Experimental Part

The experiments concerning potassium sulfate dissolution were carried out at the laboratory plant the main part of which was a vertical cylinder with the diameter of 0.12 m. The cylindrical part is finished by a conic bottom. The grid with holes is situated between the cylindrical and conic parts. The dissolution is carried out periodically, only air is supplied constantly to intensify the process. The air consumption is determined by a rotameter.

The procedure was as follows. Distilled water (2 dm\(^3\)) was loaded into the cylinder while the air compressor was switched on. After establishing the definite value of the air consumption the salt sample was loaded and stopwatch was started. In definite periods of time the samples were withdrawn and the salt concentration in the sample was determined. The experimental temperature was 293 ± 0.5 K. The fraction of K\(_2\)SO\(_4\) with the average diameter of 4·10\(^{-3}\) m was selected by sieving. The density of the solid phase was 2660 kg/m\(^3\) and saturation concentration was 107 kg/m\(^3\).

150 g of salt was loaded in the apparatus that was equal to 126 kg of salt at the concentration of 107 kg/m\(^3\). The fraction of K\(_2\)SO\(_4\) was loaded into the cylinder while the air compressor was switched on. After establishing the definite value of the air consumption the salt sample was loaded and stopwatch was started. In definite periods of time the samples were withdrawn and the salt concentration in the sample was determined. The experimental temperature was 293 ± 0.5 K. The fraction of K\(_2\)SO\(_4\) with the average diameter of 4·10\(^{-3}\) m was selected by sieving. The density of the solid phase was 2660 kg/m\(^3\) and saturation concentration was 107 kg/m\(^3\).
3. Results and Discussion

Fig. 1 represents the experimental results concerning the change of $K_2SO_4$ concentration in water during the definite time for three different values of gas consumption. The increase in gas consumption intensifies the dissolution process.

![Fig. 1. $K_2SO_4$ concentration vs. time at gas different consumptions ($m^3/s$): 2.0$ \times 10^{-3}$ (1); 5.0$ \times 10^{-3}$ (2) and 1.0$ \times 10^{-2}$ (3)](image)

It was mentioned above that dissolution is characterized by statics and kinetics. The statics is described by the material balance of Eq. (4). Let us represent this equation in the dimensionless form: both concentrations $c$ and $c_{\text{max}}$ are divided by the saturation concentration $c_{\text{sat}}$:

$$\frac{c}{c_{\text{sat}}} = \tau$$ and $$\frac{c_{\text{max}}}{c_s} = \tau_{\text{max}}$$

Apparently the relative concentrations are changed from 0 to 1 and Eq. (4) has the form:

$$\eta = 1 - \frac{1}{\tau_{\text{max}}} \cdot \tau$$

The value $\tau_{\text{max}}$ is equal to 0.7 under the experimental conditions. Fig. 2 represents the working line which is the same for all three cases.

Using the experimental data (Fig. 1) and dependence (11) we determine the values $a$ and $a_b$ according to Eq. (12). Values $a$ is 0.753 under the experimental consitions.

The difference $\left[ \phi \left( \frac{1}{a} \right) - \phi \left( \frac{\varphi}{a} \right) \right]$ as a function of time $t$ is represented by the plot (Fig. 3) depending on gas consumption.

![Fig. 2. The working line of the dissolution process](image)

![Fig. 3. $\left[ \phi \left( \frac{1}{a} \right) - \phi \left( \frac{\varphi}{a} \right) \right]$ vs. dissolution time $t$](image)

The slope angle $\gamma$ has a value of $t g \gamma = \frac{c_{\text{max}} \cdot a^2}{\rho_s \cdot d_0} \cdot \frac{2 \cdot \hat{\beta}}{H}$. Hence, $\beta = t g \gamma \cdot \frac{\rho_s \cdot d_0}{c_{\text{sat}} \cdot a^2}$.

We compare the obtained results with theoretically calculated values according to Eq. (2) on the basis of the local isotropic turbulence theory. The main value affecting the mass transfer coefficient $\beta$ under other constant parameters is a specific energy $e_0$. It is determined as follows:

$$e_0 = \frac{E}{m} = \frac{V_g \cdot \Delta \rho}{m} = \frac{V_g \cdot \rho_{\text{sat}} \cdot g \cdot H}{m}$$

where $\Delta \rho$ – pressure loss, Pa; $m$ – suspension mass, kg; $V_g$ – gas consumption, $m^3/s$; $\rho_{\text{sat}}$ – suspension density, $kg/m^3$ and $H$ – suspension height, m.
Theoretical and experimental values of the mass transfer coefficient $\beta$

<table>
<thead>
<tr>
<th>$V_g$, m/s</th>
<th>$e_0$, W/kg</th>
<th>$\beta_{theor}$, m/s</th>
<th>$\beta_{exp}$, m/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 $\times$ 10$^{-3}$</td>
<td>1.768</td>
<td>5.96 $\times$ 10$^{-3}$</td>
<td>5.8 $\times$ 10$^{-3}$</td>
</tr>
<tr>
<td>5 $\times$ 10$^{-3}$</td>
<td>4.42</td>
<td>7.49 $\times$ 10$^{-3}$</td>
<td>7.5 $\times$ 10$^{-3}$</td>
</tr>
<tr>
<td>1 $\times$ 10$^{-2}$</td>
<td>8.82</td>
<td>8.9 $\times$ 10$^{-5}$</td>
<td>8.4 $\times$ 10$^{-5}$</td>
</tr>
</tbody>
</table>

The coefficient of molecular diffusion $D$ is equal to $1.12 \times 10^{-9}$ m$^2$/s [2].

The calculated theoretical values of the mass transfer coefficient are represented in Table depending on the gas consumption. They are also compared with the experimental ones. The data given in Table show the satisfactory correlation between the experimental and theoretical values of the mass transfer coefficient $\beta$ within the range of specific energy $1.7 < e < 8.8$ kW/kg. Thus, the values calculated according to Eq. (2) may be used for calculations of the dissolution process with a satisfactory degree of reliability to determine apparatus size and dissolution time.

### 4. Conclusions

The dissolution kinetics of the solid body in the apparatus with the pneumatic agitation was investigated and the dependence of concentration on time was established.

We solved the differential kinetic equation while changing the particle diameter and process motive force.

The mass-transfer coefficients were determined during dissolution and they were compared with the theoretical values obtained on the basis of local isotropic turbulence.

### References


АНАЛІЗ КІНЕТИКИ РОЗЧИНЕННЯ НА ОСНОВІ ТЕОРІЇ ЛОКАЛЬНОЇ ІЗОТРОПНОЇ ТУРБУЛЕНТНОСТІ

Анотація. Досліджено кінетику розчинення сульфату калію у воді при пневматичному перемішуванні системи. Визначено коефіцієнти масовіддачі на основі експериментальних досліджень та теоретичного аналізу різних теорій локальної ізотропної турбулентності. Показано адекватність розрахунків теоретичних та експериментальних даних, що дає можливість теоретичного визначення коефіцієнтів масовіддачі з метою розрахунку даного процесу.

Ключові слова: розчинення, кінетика розчинення, коефіцієнт масовіддачі, інтенсифікація.