Evaluation of Adding Carbon Tetrachloride as Propulsion to the Thermal Cracking Reactor due to the Amount of Formed Coke in Different Coil Outlet Temperatures (COT)

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Abstract

Nowadays, polyvinyl chloride (PVC) was the most usual plastic. Thereby, to make polyvinyl chloride its monomer must be produced firstly that was called vinyl chloride (VCM). It was severely endothermic reaction that was done in an ethylene dichloride ethylene dichloride thermal cracking reactor within temperature range of 680-758 K and pressure range of 2500000 Pascal, thus this cracking reaction changed into hydrochloric acid and VCM. In production unit, monomeric chloride had the main and principal role as the core of the process of thermal cracking that occurs in the furnace. Increasing of wall temperature cause to boil gas mixture and causing pyrolysis reactions. Regarding to the simulation results showed that number of pyrolysis produced composition have maximum concentration in the length of reactor that illustrated these compositions participated in secondary reactions. Furthermore, by increasing the amount of coil outlet temperatures, the amount of formed coke will be increased. If Carbon tetrachloride considered as the chlorine radicals, it has an important role as the motivator in the cracking procedures, radicals causing an enhancement in VCM production.

Keywords: Carbon tetrachloride; Conversion factor; Coke; Coil outlet temperatures; Thermal cracking

Introduction

Design and Simulation of thermal cracking reactor because of the chemical reactions in the processes and heat transfer in the furnace are very complex. Therefore, this process from the chemical reactions engineering perspective and from the perspective of transfer phenomena is highly regarded. Computational Fluid Dynamics (CFD) tools help us to investigate the development of detailed models of these reactors and by understanding the basic principles, the optimized design of reactor and by improving the cost and time performance of this equipment in a virtual laboratory can be done. In this research, investigate the proposed model of EDC cracking process in an industrial furnace and by operational data from a petrochemical unit to qualify the model and remove the reductions. To begin, by previous researches that worked on this specific occasion were collected. Then they are studied and investigated. Moreover, the reactor mathematical models were provided and solved by numerical software. Thereby, the operation of the reactor at different operating conditions investigated and sensitivity analysis of effective parameters will be done. As a result, this work could design new reactors or increasing the optimization capacity of existing reactors.

Molecular cracking

Molecular cracking was a process that used in the petrochemical industry and to reduce their hydrocarbons molecular weight by breaking their bonds. This process is the main principal of converting crude oil to useful fuels such as gasoline, diesel, jet fuel and kerosene. Thermal cracking, catalytic cracking, hydrocracking and cracking by water vapour were the most useful cracking processes in the industry. This process was done in the high temperature and pressure and without presence of catalyst or in the low temperature and pressure with presence of catalyst.

Geometry of thermal cracking reactor

In EDC thermal cracking process, coils containing process fluid, inside a box called furnace, are positioned horizontally. Coils placement inside the furnace shown in Figure 1 [1]. That positioned on the on both sides of the wall furnace, by burning gases of these furnaces the heat produced, that the heat transferred to the coils by convection or radiation model (Table 1). The number of intended coils to simulate 11 rows of pipes (Inconel 600) with overall length of 95 meters and length of each row tubing is 13.5 meters. Inner diameter of coils is 0.134 meter the distance between centers of the two tubes is 0.32 meter (Figures 2 and 3).

Mass balance equation in the software proposed in form of equation 1

$$\frac{\partial (\rho u)}{\partial t} + \frac{\partial (\rho u^2)}{\partial x} = \frac{\partial J}{\partial x} + R + S$$  

Where:

- $\rho$: Density
- $t$: time
- $u$: Potential energy
- $J$: diffusion term
- $R$: input heat to the reactor
- $S$: heat at point $i$

Phrase $J$ reflects is the term infiltration, which is not used in this simulation [2,3]. The energy equation in the Fluent software as follows:

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Figure 1: Placement of coil in the furnace.

Figure 2: Schematic of one row of the coil in the software.

Figure 3: Prospective a row of coil.

\[ s_i = \text{input thermal of chemical reactions} \]
\[ S_A = \text{thermal of chemical reactions} \]
\[ K_{\text{eff}} = \text{the effective conductivity} \]
\[ E = \text{total energy} \]

\[ K_{\text{eff}} + K = K_{\text{eff}} \]

is the effective conductivity, \( K \) turbulent thermal conductivity, according to the model used is the perturbation. The first three terms on the right side of the equation as an expression of directed energy transfer, distribution of components and dispersion of viscosity. \( S_A \) is thermal of chemical reactions [4-6]. \( E \) is expressed as follows:

\[ E = h - \frac{p}{\rho} + \frac{u_i^2}{2} \]  

(3)

\[ u_i = \text{input Potential energy} \]
\[ p = \text{pressure} \]
\[ \rho = \text{Density} \]
\[ E = \text{total energy} \]
\[ u_i = \text{input Potential energy} \]
\[ h = \text{input heat} \]

Mass continuity equation in FLUENT software is in form of eqn 4:

\[ \frac{\partial \rho}{\partial t} + \frac{\partial \rho u_i}{\partial x_j} = 0 \]  

(4)

Above equation was the general form of the mass balance equation for compressible and non-compressible flows.

\[ \frac{\partial}{\partial t} \left( \rho u_i \right) + \frac{\partial}{\partial x_j} \left( \rho u_i u_j \right) = -\frac{\partial}{\partial x_j} \left( \rho g_i \right) + \rho \frac{\partial \tau_{ij}}{\partial x_j} + F_i \]  

(5)

In the above equation, \( P \) static pressure, \( \tau \) tensor stress, \( \rho g \) and \( F \) physical forces of gravity and physical external factors are in \( i \) direction. Tensor stress is as follows:

\[ \tau_{ij} = \left[ \mu \left( \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} \right) \right] - \frac{2}{3} \mu \frac{\partial u_i}{\partial x_j} \delta_{ij} \]  

(6)

\( \mu \) the molecular viscosity and the second semester is the influence of volumetric expansion [3].

**Standard k-\( \varepsilon \) model**

This was a comprehensive model that account as the simplest turbulence model. Two educational turbulent models enables solving the two transfer equations that separately determined turbulent length and time scale. Standard k-\( \varepsilon \) model is useful for powerful engineering calculations. Powerful of this equation, economical and acceptable accuracy for a wide range of turbulent flow as a result of its popularity in industrial flow and heat transfer simulation. This model is a semi-empirical model. Standard k-\( \varepsilon \) model, was a model based on transfer equations for kinetic turbulent energy and dissipation rate (\( \varepsilon \)). In achieving the standard k-\( \varepsilon \) model assumes that the fully turbulent flow and molecular viscosity effects have been ignored. Because of the strengths and weaknesses of the equation to improve its performance, k-\( \varepsilon \) realizable and k-\( \varepsilon \) RNG model was introduced. K and \( \varepsilon \) is obtained by solving simultaneously the following two equations:
Table 1: Energy of EDC thermal cracking reaction [2].

<table>
<thead>
<tr>
<th>Reactions</th>
<th>A([cm^3/mole]*n's^-1)]</th>
<th>n</th>
<th>E (kJ/mole)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radical reaction chain initiation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EDC + R1 → HCl + R2</td>
<td>5.9 × 1015</td>
<td>1</td>
<td>342</td>
</tr>
<tr>
<td>CCl4 + R1 → R1</td>
<td>2.2 × 1012</td>
<td>1</td>
<td>230</td>
</tr>
<tr>
<td>Radical reaction chain propagation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EDC + R1 → HCl + R2</td>
<td>1.3 × 1013</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td>EDC + R1 → VM + R3</td>
<td>1.2 × 1013</td>
<td>2</td>
<td>34</td>
</tr>
<tr>
<td>EDC + R1 → EC + R2</td>
<td>1.0 × 1012</td>
<td>2</td>
<td>42</td>
</tr>
<tr>
<td>EDC + R1 → 1,1,2+R4</td>
<td>5.0 × 1011</td>
<td>2</td>
<td>45</td>
</tr>
<tr>
<td>EDC + R1 → 1,1,1,2+R5</td>
<td>2.0 × 1011</td>
<td>2</td>
<td>48</td>
</tr>
<tr>
<td>EDC + R1 → 1,1,1,2+R6</td>
<td>1.0 × 1011</td>
<td>2</td>
<td>56</td>
</tr>
<tr>
<td>EDC + R1 → CHCl + R5</td>
<td>1.0 × 1012</td>
<td>2</td>
<td>63</td>
</tr>
<tr>
<td>VCM + R1 → R1</td>
<td>9.1 × 1010</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>VCM + R1 → HCl + R1</td>
<td>1.2 × 1014</td>
<td>2</td>
<td>56</td>
</tr>
<tr>
<td>VCM + R1 → CP + R1</td>
<td>5.0 × 1011</td>
<td>2</td>
<td>31</td>
</tr>
<tr>
<td>VCM + R1 → C2H2 + R1</td>
<td>2.0 × 1010</td>
<td>2</td>
<td>30</td>
</tr>
<tr>
<td>VCM + R1 → EC + R2</td>
<td>3.0 × 1011</td>
<td>2</td>
<td>61</td>
</tr>
<tr>
<td>R1 → VCM + R5</td>
<td>2.1 × 1014</td>
<td>1</td>
<td>84</td>
</tr>
<tr>
<td>R1 → C2H2 + R5</td>
<td>5.0 × 1014</td>
<td>1</td>
<td>90</td>
</tr>
<tr>
<td>R1 → Di + R1</td>
<td>2.0 × 1013</td>
<td>1</td>
<td>70</td>
</tr>
<tr>
<td>R1 → Tri + R1</td>
<td>2.5 × 1013</td>
<td>1</td>
<td>70</td>
</tr>
<tr>
<td>2C2H + R1 → C2H2 + R1</td>
<td>1.0 × 1014</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td>EC + R1 → HCl + R5</td>
<td>1.7 × 1013</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>1,1+R1 → HCl + R5</td>
<td>1.2 × 1013</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>1,1,2+R2 → HCl + R5</td>
<td>1.7 × 1013</td>
<td>1</td>
<td>15</td>
</tr>
<tr>
<td>1,1,1,2+R4 → HCl + R5</td>
<td>1.7 × 1013</td>
<td>2</td>
<td>17</td>
</tr>
<tr>
<td>CHCl + R1 → HCl + R5</td>
<td>1.6 × 1013</td>
<td>2</td>
<td>14</td>
</tr>
<tr>
<td>CCl4 + R1 → HCl + R5</td>
<td>5.0 × 1011</td>
<td>2</td>
<td>33</td>
</tr>
<tr>
<td>CCl4 + R4 → 1,1,2+R6</td>
<td>1.0 × 1012</td>
<td>2</td>
<td>33</td>
</tr>
<tr>
<td>CCl4 + R4 → 1,1,1,2+R7</td>
<td>5.0 × 1012</td>
<td>2</td>
<td>33</td>
</tr>
<tr>
<td>Radical reaction chain termination</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R1 → VCM + HCl</td>
<td>1.0 × 1013</td>
<td>2</td>
<td>13</td>
</tr>
<tr>
<td>R1 + Di → HCl + R5</td>
<td>1.0 × 1013</td>
<td>2</td>
<td>12</td>
</tr>
<tr>
<td>R1 + Di → CCl4 + R5</td>
<td>1.0 × 1013</td>
<td>2</td>
<td>13</td>
</tr>
<tr>
<td>C2H + 2R1 → 2C2H2 + R1</td>
<td>1.6 × 1014</td>
<td>2</td>
<td>70</td>
</tr>
</tbody>
</table>

**RNG k-ε model**

This model is based on statistical methods, this model is the same as the standard k-ε model but with the following modifications:
1. An additional term (ε) has been added that significantly increases accuracy for flow to layer velocity.
2. The effect of the rotational movement (vortex motion) included in turbulences and increases accuracy for rotary flow.
3. An analytical formula for turbulent Prantel numbers are provided, as standard k-ε model considers a fixed amount.
4. While the standard k-ε model is the model for high Reynolds numbers, the theory of RNG for analysis of a differential equation for the turbulent viscosity can be provided that it considered the effects of low Reynolds. Effective use of this feature depends on proper behavior of the region near the wall of furnace. k-ε realizable model.

This model has limitations on Reynolds stresses that match with the physics of turbulent flows. This is possibly not in the standard k-ε model and not in the k-ε RNG model. In places where a large curvature in flow lines, vortex and rotation of the screw. K-ε RNG and k-ε realizable models have significant progress towards standard k-ε model. Preliminary studies show that the k-ε realizable model between k-ε models have best performance for confirmation numbers of complex flows and segregated flows. For all of this cases the performance of this model is better than the standard k-ε model. As it was observed that in the standard k-ε model, turbulent viscosity achieved the following equation:

$$\mu_t = \rho C_{\mu} \frac{K^2}{\varepsilon}$$  \hspace{1cm} (10)

That $C_{\mu}$ is constant, but in the k-ε realizable $C_{\mu}$ achieved by the following equation:

$$C_{\mu} = \frac{1}{A_{\mu} + A_{\mu} \frac{KU}{\varepsilon}}$$ \hspace{1cm} (11)

$$U^* = \sqrt{S_1 S_2 S_3} \Omega^*$$ \hspace{1cm} (12)

$$\Omega^* = \Omega - 2\varepsilon \varepsilon \Omega$$ \hspace{1cm} (13)

$$\Omega^* = \Omega^* - \xi \varepsilon \Omega$$ \hspace{1cm} (14)

$\Omega^*$ Average rate of tensor with rotational speed is $\omega_i$ constants are defined below.

$$A_{\mu} = 4/9 A_{\mu} = \sqrt{6 \cos \phi}$$ \hspace{1cm} (15)

$$S_{ij} = \frac{1}{2} \frac{\partial U_j}{\partial x_i} + \frac{\partial U_i}{\partial x_j}$$ \hspace{1cm} (16)

As it’s be seen $C_{\mu}$ placed a function of the average strain, rotational speeds, the angle speed of rotation systems, and turbulent fields are (k and ε) (Table 2).

**Wall of furnace**

Profile of heat was applied in the walls. A total of eleven wall was placed a function of the average strain, rotational speeds, the angle speed of rotation systems, and turbulent fields are (k and ε) (Table 2).
The transfer coefficient (W/m·K) and the other base profile points due to eq.17 and eq.18 multiply in a constant parameter to obtain new Profile points.

\[
\text{Weight factor}(x) = \frac{\text{Temperature}(x) \cdot \text{Temperature}(l = 0)}{\text{Temperature}(l = 150) \cdot \text{Temperature}(l = 0)}
\]

(17)

\[
T(x) = \text{Weight factor}(x) \times \frac{T(l = 150)_{\text{desired}}}{774.1} \times T_{\text{base}}
\]

(18)

The results of these calculations can be seen in Figure 5. COT different temperature profiles have been obtained. Figure 6 indicates the EDC conversion factor in different COTs as the figure shows at higher conversion factor it will be higher in the COTs. But according to the Figure 7 if the amount of COT is higher, the amount of formed coke also will be higher. So by regarding the simulation results it can be said that for optimum economic performance of furnace, the temperature profile of the fluid is very important.

**Investigating of adding carbon tetrachloride as propulsion**

By adding this compound to the feed in addition to increase conversion factor, it has other effects on the process. Due to increasing number of chlorine radicals in the process, number of unwanted compounds and Acetylene production and as a result coke production has been increased. Creating unwanted compounds in the next stages, in addition to VCM treating problem, also an important loss costly. As shown in Figure 8 conversion factor increases with increasing carbon tetrachloride, on the other hand in the Figure 9 formed coke, of the increases in carbon tetrachloride. So regarding to the simulation results it can be said that for optimum economic performance of furnace, the temperature profile of the fluid is very important. Therefore, optimize the amount of added carbon tetrachloride to the feed for economic performance of thermal cracking unit is very important. In this amount of propulsion conversion factor increases as an allowable limit and reactor performance due to forming coke and unwanted compounds are economical. It can be seen in the Figure 5.

**Conclusion**

K-ε realizable model is the most appropriate model for calculation because of these reasons: [7,8]
This version contains a new formula for turbulence viscosity.

A modified transferring equation for scattering rate, ε, can be considered. Therefore, this model is an exact equation for fluctuations transferring has been made.

In this research, effect of added carbon tetrachloride as a propulsion has been investigated. According to result of this research and other issues, if the amount of this compound does not reach more than 100 ppm, conversion factor will be increased. If this amount will be more than 100 ppm it’s not economically suggested.

References