Migration and Transformation of Vanadium and Nickel in High Sulfur Petroleum Coke during Gasification Processes

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Keywords: migration, sequential extraction, gasification, nickel, high sulfur petcoke, vanadium

Abstract:

The volatilization characteristics and occurrence forms of V and Ni in petroleum coke (petcoke) were investigated during steam (H\textsubscript{2}O) and carbon dioxide (CO\textsubscript{2}) gasification on a fixed bed reactor at 800\textdegree\textendash1100 °C. The Tessier sequential chemical extraction procedure was employed to determine the different forms of V and Ni. The results showed their volatilities were not dependent on the gasification atmosphere, but rather relied mainly on the reaction temperature. The CO\textsubscript{2} atmosphere accelerated the conversion of organic-bound nickel to residual form at low temperature and promoted Fe-Mn oxides formation at high temperature. However, the H\textsubscript{2}O atmosphere was conducive to form vanadium bound to Fe-Mn oxides and promoted the decomposition of residual forms. In addition, the thermodynamic equilibrium calculations showed the volatilization of Ni mainly released Ni\textsubscript{S}\textsubscript{2} between 800\textdegree\textendash1100 °C. The H\textsubscript{2}O atmosphere was favorable to generate the more stable Ni\textsubscript{x}S\textsubscript{y} compound, thereby suppressing the volatilization of Ni, while the presence of CO\textsubscript{2} led to an increase in residual V and decrease of Fe-Mn oxides. The V and Ni mainly caused erosion problems under the CO\textsubscript{2} atmosphere while the fouling and slagging obviously increased under the H\textsubscript{2}O atmosphere with impacts gradually weakened with the increase of temperature.

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Abstract: The volatilization characteristics and occurrence forms of V and Ni in petroleum coke (petcoke) were investigated during steam (H₂O) and carbon dioxide (CO₂) gasification on a fixed bed reactor at 800–1100 °C. The Tessier sequential chemical extraction procedure was employed to determine the different forms of V and Ni. The results showed their volatilities were not dependent on the gasification atmosphere, but rather relied mainly on the reaction temperature. The CO₂ atmosphere accelerated the conversion of organic-bound nickel to residual form at low temperature and promoted Fe-Mn oxides formation at high temperature. However, the H₂O atmosphere was conducive to form vanadium bound to Fe-Mn oxides and promoted the decomposition of residual forms. In addition, the thermodynamic equilibrium calculations showed the volatilization of Ni mainly released Ni₃S₂ between 800–1100 °C. The H₂O atmosphere was favorable to generate the more stable NiₓSᵧ compound, thereby suppressing the volatilization of Ni, while the presence of CO₂ led to an increase in residual V and decrease of Fe-Mn oxides. The V and Ni mainly caused erosion problems under the CO₂ atmosphere while the fouling and slagging obviously increased under the H₂O atmosphere with impacts gradually weakened with the increase of temperature.

Keywords: high sulfur petcoke; vanadium; nickel; gasification; sequential extraction; migration

1. Introduction

The shortage of energy resources and the improvement of petroleum processing capacity have led to a continuous increase in the production of petcoke [1–3]. Petcoke is mainly used in the electrolytic aluminum industry and boilers as a combustion fuel, however it releases massive amounts of polluting gases such as SO₂ under combustion conditions, resulting in serious environmental problems [4]. Petcoke has high calorific value, low ash and low price qualities, which endow petcoke with the potential to be a gasification feedstock to produce synthetic gas and recycle sulfur through the Claus process. Therefore, it is an efficient and clean utilization way, which has broad prospects for making waste profitable [5,6]. Unfortunately, petcoke is enriched with Ni and V concentrations of more than 1000 ppm, resulting in serious fouling, slagging and erosion problems of gasification equipment and pipes [7]. V₂O₅ can react with refractory and metal oxides [8], exposing the surface of the metal to an oxidizing atmosphere and accelerating its oxidation rate. High concentrations of V, Ni and sulfur in petcoke can cause severe pollution, including erosion and corrosion to the furnace and equipment, during gasification [9]. Therefore, a thorough understanding of the chemical speciation migration of V and Ni in the process of gasification and the complex chemical reactions that may occur with...
gas reactants and mineral components is important to predict and control the fouling, slagging and erosion behavior during the process of gasification.

There are three gasification technologies, including fixed bed, fluidized bed, and entrained flow bed, of which the entrained flow bed appears to be a promising gasification technology with the advantages of high output, high thermal conversion efficiency because of its higher operating temperatures. However, the high temperature also accelerates the fouling, slagging and erosion problems caused by vanadium and nickel in the petcoke. Park et al. [10] investigated slagging of petroleum coke ash. The main components of petcoke under gasification conditions were vanadium, nickel and iron. Among them, vanadium existed in the form of \( \text{V}_2\text{O}_3 \) with a melting point of up to 1940 °C. \( \text{V}_2\text{O}_3 \) in petcoke ash remained solid at the typical gasification temperature and the solid \( \text{V}_2\text{O}_3 \) caused the slag viscosity to increase. For the pet coke with high ash content, the effect of \( \text{V}_2\text{O}_3 \) on slag viscosity must be taken into consideration when the minimum gasification temperature was estimated. Nakano et al. [11] studied a synthetic slag and found that \( \text{V}_2\text{O}_3 \) could form solid solutions with \( \text{Al}_2\text{O}_3 \) and \( \text{Fe}_2\text{O}_3 \) to reduce the slag viscosity. Duchesne et al. [12] examined the viscosity characteristics of coal and petcoke blended ash. Petcoke ash was mainly composed of V and Ni, and the spinel formed by vanadium could increase the ash viscosity. Mahapatra et al. [13] measured the release of vanadium and nickel during the pyrolysis process, reporting that vanadium and nickel tended to accumulate in the coke compared to fly ash. Wang et al. [7] investigated that the formation of ash and slag was closely related to the conversion of vanadium, nickel, and sulfur during the gasification process. Brüggemann et al. [14] also observed that large amounts of gaseous nickel-containing compounds were formed in the synthesis gas. Therefore, the above problems can be relieved by using a relatively low reaction temperature.

The investigation of the occurrence and migration behavior of V and Ni has been studied by many researchers. Querol et al. and Dai, et al. [15,16] expressed that vanadium existed in the form of organic matter and aluminosilicate minerals in coal. Brüggemann et al. [14] indicated that nickel was transformed into \( \text{Ni}_x\text{S}_y \) and elemental Ni under high \( \text{H}_2\text{S} \) and low \( \text{H}_2\text{S} \) partial pressure, which was calculated from the thermodynamics of the conversion of Ni during heavy oil gasification. Wang et al. [17] found that \( \text{V}_2\text{O}_5 \) reacts with \( \text{CaO} \) to form \( \text{Ca}_3\text{V}_2\text{O}_8 \) when blended with coal ash in a reducing atmosphere, and the decomposition of \( \text{Ca}_3\text{V}_2\text{O}_8 \) could react with \( \text{Fe}_2\text{O}_3 \) to form \( \text{FeV}_2\text{O}_4 \) when the temperature was over 1300 °C. Li et al. [18] reported that the vanadium reacted with the mineral component to form a new thermal stable compound when the gasification temperature was above 1300 °C. However, a detailed systematic research on primary speciation of V and Ni in petcoke and its release and migration characteristics during gasification has not been accomplished.

Therefore, in this study, the behavior of V and Ni in the \( \text{CO}_2 \) and \( \text{H}_2\text{O} \) gasification of petcoke was studied by fixed bed gasification experiments. The Tessier sequential chemical extraction procedure and ICP-MS were used to obtain the forms of V and Ni in the gasification product under different gasification conditions. Furthermore, the release and transformation mechanism of V and Ni were elaborated deeply. Moreover, thermodynamic equilibrium calculations, the use of the FACTSAGE software, were carried out to study the reaction mechanism of V and Ni with the main associated mineral elements during gasification. Finally, a detailed assessment on the pernicious effects on gasification equipment by the mineral elements V and Ni was gained.

2. Experiments

2.1. Petcoke Samples

The petcoke samples were obtained from the Wuhan branch of Sinopec, ground and sieved through 100 mesh sieves. Before the gasification experiment, the samples were dried at 80 °C in advance in a vacuum drying oven for 24 h. The petcoke ash was prepared at 805 °C for 2 h according to the Chinese standard procedures (GB/T 1574-2007). Table 1 shows the ultimate and proximate analysis of the samples. Table 2 lists the ash compositions (by XRF) of the petcoke.
Table 1. Proximate and ultimate analysis of petcoke samples (wt %).

<table>
<thead>
<tr>
<th></th>
<th>( M_{\text{ad}} )</th>
<th>( A_{\text{ad}} )</th>
<th>( V_{\text{ad}} )</th>
<th>( F_{\text{ad}} )</th>
<th>( C_{\text{ad}} )</th>
<th>( H_{\text{ad}} )</th>
<th>( N_{\text{ad}} )</th>
<th>( S_{\text{ad}} )</th>
<th>( O_{\text{ad}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.46</td>
<td>0.50</td>
<td>8.56</td>
<td>89.48</td>
<td>85.43</td>
<td>3.63</td>
<td>1.81</td>
<td>2.01</td>
<td>7.12</td>
</tr>
</tbody>
</table>

\( M_{\text{ad}} \): the proximate and ultimate analysis of the samples are on the air-dry basis; *: the ultimate analysis of O is quantified by the subtraction method.

Table 2. Ash composition of petcoke samples (wt %).

<table>
<thead>
<tr>
<th></th>
<th>NiO</th>
<th>( V_2O_5 )</th>
<th>( Fe_2O_3 )</th>
<th>SiO(_2)</th>
<th>SO(_3)</th>
<th>Al(_2)O(_3)</th>
<th>TiO(_2)</th>
<th>CaO</th>
<th>Na(_2)O</th>
<th>MnO</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>20.11</td>
<td>14.88</td>
<td>9.41</td>
<td>18.18</td>
<td>3.87</td>
<td>13.59</td>
<td>4.29</td>
<td>14.54</td>
<td>4.29</td>
<td>0.17</td>
</tr>
</tbody>
</table>

2.2. Fixed Bed Gasification Apparatus and Experimental Procedures

Our laboratory-scale reactor system (Figure 1) contains a steam generator and a fixed-bed reactor that is heated by an electrical furnace. The total length of the quartz tube is 500 mm and the inner diameter is 40 mm. The quartz tube is heated by an electrothermal furnace, which can reach a maximum temperature of 1100 °C under control by a temperature controller. The steam generator is mainly composed of an electrical heating furnace and a high-precision syringe pump which precisely controls the water injection (0.736 mL/min of deionized water provides 1.00 L/min steam at ambient temperature). The deionized water is evaporated in the preheating furnace, and then carried by the carrier gas of \( \text{N}_2 \). The mixture gas enters the quartz tube through a heat insulating pipeline wrapped with a heating tape.

![Diagram of the fixed bed gasification system.](image)

In the petcoke gasification experiments, a \( \text{N}_2 \) flow of 500 mL/min was maintained to remove air from the system. When the specified temperature was reached, the atmosphere was switched to carbon dioxide gas or steam carried by nitrogen. The flow rate of \( \text{CO}_2 \) was kept at 500 mL/min for all \( \text{CO}_2 \) gasification experiments, while that of \( \text{N}_2 \) was kept at 150 mL/min and \( \text{H}_2\text{O} \) at 350 mL/min for all \( \text{H}_2\text{O} \) gasification experiments, respectively. Meanwhile, basket with 1 g samples was rapidly placed in the quartz mesh in the reaction zone. The reaction time was 30 min. After gasification, the basket was pulled to the up and cold position and rapidly cooled to room temperature by \( \text{N}_2 \). The gasification chars were collected, weighted, and subsequently analyzed.

2.3. Sequential Chemical Extraction

Sequential chemical extraction was an effective way to quantify the chemical forms of trace elements [19–21]. In this research, five chemical forms of V and Ni in petcoke and char after gasification were extracted by the Tessier sequential extraction procedure [22]. In the extraction procedures [23],
the Ni and V were extracted and partitioned into five forms: (1) exchangeable, extracted by MgCl$_2$ (1 mol/L); (2) bound to carbonates, extracted by CH$_3$COONa (1 mol/L); (3) bound to Fe-Mn oxides, extracted by NH$_2$OH·HCl (0.04 mol/L) which dissolved in CH$_3$COOH (25%, v/v); 4) bound to organic matter, extracted by HNO$_3$ (0.02 mol/L) and H$_2$O$_2$ (30%, v/v); After four extraction steps, Ni and V still remaining in the residue were labeled as residual forms, which might be present in the insoluble organic matrix or occur in minerals such as aluminosilicates, titanium dioxide, zircon, sulphides and so on [18,24].

2.4. Calculation of V and Ni Volatility

The V and Ni contents in petcoke and its gasification char and their extracted solutions were measured by inductively coupled plasma mass spectrometry (ICP-MS). V and Ni volatility is determined by the equation:

$$R_v(\%) = \left(1 - \frac{C_1 m_1}{C_0 m_0}\right) \times 100\% = \left(1 - \frac{C_1 Ry}{C_0}\right) \times 100\%$$

Char yield after petcoke gasification is calculated by $R_y$, which is assessed as follows:

$$R_y(\%) = \left(\frac{m_1}{m_0}\right) \times 100\%$$

where $R_v$ is the volatility of V or Ni, %; $R_y$ is the yield of gasification char, %; $C_1$, the concentration of the V or Ni in gasification char, ug/g; $C_0$, the concentration of V or Ni in petcoke, ug/g; $m_1$, the weight of gasification char, g; $m_0$, the weight of petcoke, g.

2.5. Thermodynamic Equilibrium Calculations

The FACTSAGE program is used to facilitate our explanation of the transformation mechanisms of V and Ni during gasification and verify the experimental results. FACTSAGE [25] software is based on the Gibbs energy minimization method. The Equilib module utilized with the FactPS and FToxid databases can detect the concentration of chemical species when specific elements or compounds react to equilibrium. The precision of the software can meet the demands of simulation. In this research, modeling employed main trace elements on behalf of the main composition of petcoke. Simulation calculations were carried under the temperature range between 100 °C to 1500 °C at normal atmospheric pressure. The quantity of petcoke feedstock to be fed to the gasifier was 1 kg and the CO$_2$ or H$_2$O was 1.5 kg. It was noticeable that the quantity of gasification agent and mineral elements was in units of g/kg-petcoke in the gasification experiment. However, the unit (g/kg-petcoke) was modified to mol/kg-petcoke for the purpose of calculation. The input information is listed in Table 3. There is an assumption that all the sulfur in petcoke is presented in the vapor phase as H$_2$S (g) initially because all of gasifiers produce H$_2$S (g) [18,26].

<table>
<thead>
<tr>
<th>Reactants</th>
<th>mol/kg-Raw Petcoke</th>
<th>Reactants</th>
<th>mol/kg-Raw Petcoke</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>71.19</td>
<td>NiO</td>
<td>$1.31 \times 10^{-2}$</td>
</tr>
<tr>
<td>H$_2$</td>
<td>17.52</td>
<td>V$_2$O$_5$</td>
<td>$0.40 \times 10^{-2}$</td>
</tr>
<tr>
<td>N$_2$</td>
<td>0.65</td>
<td>Fe$_2$O$_3$</td>
<td>$0.29 \times 10^{-2}$</td>
</tr>
<tr>
<td>H$_2$S</td>
<td>0.63</td>
<td>SiO$_2$</td>
<td>$1.49 \times 10^{-2}$</td>
</tr>
<tr>
<td>CO$_2$*1</td>
<td>34.00</td>
<td>Al$_2$O$_3$</td>
<td>$0.65 \times 10^{-2}$</td>
</tr>
<tr>
<td>H$_2$O*2</td>
<td>83.33</td>
<td>TiO$_2$</td>
<td>$0.06 \times 10^{-2}$</td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>CaO</td>
<td>$1.27 \times 10^{-2}$</td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>Na$_2$O</td>
<td>$0.34 \times 10^{-2}$</td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>MnO</td>
<td>$0.01 \times 10^{-2}$</td>
</tr>
</tbody>
</table>

*1 only in the CO$_2$ gasification, *2 only in the H$_2$O gasification.
3. Results and Discussion

3.1. The Char Yield from Two Types of Gasification Atmospheres

The char yield under CO$_2$ and H$_2$O gasification conditions are shown in Figure 2. The yield tends to decrease with increasing gasification temperatures. The char reduces from 89.8% (800 °C) to 64.9% (1100 °C) in CO$_2$ gasification while the char yield declines from 80.8% (800 °C) to 29.0% (1100 °C) in H$_2$O gasification. Apparently, the atmosphere plays an important role for the yield of char and petcoke has high gasification reactivity under H$_2$O atmosphere. From the reduction trend of char yield, it can be seen there are two stages in the CO$_2$ and H$_2$O gasification. In stage 1 (before 1000 °C), the reduction trend of char yield in H$_2$O atmosphere is dramatically higher than that in CO$_2$ atmosphere. It is interesting that the reduction trend of char yield in the CO$_2$ and H$_2$O gasification in stage 2 (after 1000 °C) is opposite to stage 1 because CO$_2$ molecules can cross the reaction energy barrier leading to an increase of the char consumption yield while the H$_2$O gasification almost complete at this temperature.

![Figure 2](image-url)

**Figure 2.** Char yield (Ry%) during petcoke gasification from 800 °C to 1100 °C.

3.2. Occurrence Modes of Nickel in Petcoke and Its Gasification Char

The migration of occurrence forms of nickel during CO$_2$ and H$_2$O gasification is shown in Figure 3a,b. A few Fe-Mn oxides with a proportion of 7.9% is in the raw petcoke while the content of nickel exists in exchangeable and carbonates are tiny amounts of 0.7% and 0.3%. The majority of nickel is in organic (62.7%) and residual form (28.4%). After gasification, significant change of the different forms of nickel occurs in the gasification char.

![Figure 3](image-url)

**Figure 3.** (a) Occurrence modes of Ni in raw petcoke and char in CO$_2$ atmosphere; (b) Occurrence modes of Ni in raw petcoke and char in H$_2$O atmosphere.
For the CO\textsubscript{2} gasification, as shown in Figure 3a, when the gasification temperature increases to 800 °C, most of the organic-bound nickel is converted to the residual form. The residual form of nickel increases from 28.4% to 79.7% while the content of organic-bound sharply reduces from 62.7% to 9.7%. The Ni bound to Fe-Mn oxides is basically constant. As the gasification temperature increases from 800 °C to 900 °C, the residual form of nickel begins to decompose and mainly transforms into organic and Fe-Mn oxides. At 1000 °C, the fraction of the organic-bound nickel also transforms into Fe-Mn oxides which reaches a maximum of 14.9%. When the temperature reaches 1100 °C, organic and Fe-Mn oxides are significantly reduced and converted into the residual form of nickel. The content of the residual form of nickel reaches 79.4%, which is basically consistent with the content in the residue of nickel form at 800 °C. We note that comparing the char samples at 800 °C and 1100 °C, it can be found that the content of nickel affiliated with the exchangeable, carbonates, Fe-Mn oxides and residual forms are basically unchanged. After gasification at 800 °C, Ni associated with organic matter sharply drops 53.0% while the residual form rises acutely (51.3%) and very low level of volatile Ni (1.4%) emerges. At 1100 °C, as residual form remains unchanged, Ni affiliated with organic matter further decreases 6.7% and converts into volatile Ni. In this sense, from room temperature to 1100 °C, there are 51.3% and 6.7% of organic matter Ni converted into residual form and volatile form, respectively. It can be deduced that the organic-bound nickel prefers to form residual fraction rather than volatilize.

For the H\textsubscript{2}O gasification, as shown in Figure 3b, a large amount of organic-bound nickel decomposes from 62.7% to 28.7% and is mainly converted into the residual form (from 28.7% to 57.3%), with a small part converted into Fe-Mn oxides at 800 °C. The residual form of nickel decreases by 13.1% at 900 °C and is mainly converted into organic and Fe-Mn oxides, while the transformation behavior from 900 °C to 1000 °C shows an opposite trend. When the temperature reaches 1100 °C, organic and Fe-Mn oxides completely decompose and convert into the residual form.

In general, the gasification temperature mainly affects the transformation of the organic-bound nickel and the residual form nickel, accompanied by a small amount of other forms. Comparing the CO\textsubscript{2} and H\textsubscript{2}O gasification processes, the trends of the different forms are roughly the same, but the contents differs greatly. At 800–900 °C, the new generated residual form nickel in CO\textsubscript{2} atmosphere is nearly three times higher than that of H\textsubscript{2}O atmosphere, indicating that CO\textsubscript{2} atmosphere is conducive to the conversion of the organic-bound nickel to the residual form at low temperature. At 1000–1100 °C, the organic-bound and the residual form nickel in the CO\textsubscript{2} atmosphere are lower than in the H\textsubscript{2}O atmosphere, indicating that the CO\textsubscript{2} atmosphere promotes the formation of Fe-Mn oxides at high temperature. In both atmospheres, the organic-bound nickel gradually decreases with temperature, but the amount in the H\textsubscript{2}O one is significantly higher than that in the CO\textsubscript{2} atmosphere, indicating that the CO\textsubscript{2} gasification are favorable to the decomposition of organic-bound nickel. The trends of the residual form Fe-Mn oxides are also generally the same. The main difference is that the peak of the content of Fe-Mn oxides in H\textsubscript{2}O atmosphere is advanced to 900 °C, which is probably due to the relatively low activity of CO\textsubscript{2} and the decomposition of bound to Fe-Mn oxides requires higher energy to carry out, leading to the transformation of most of Fe-Mn oxides into the residual form of nickel at 1000 °C under CO\textsubscript{2} atmosphere.

As the temperature increases, the volatilization characteristic of nickel gradually increases during H\textsubscript{2}O and CO\textsubscript{2} gasification, illustrated in Figure 4. It is remarkable that H\textsubscript{2}O and CO\textsubscript{2} gasification display similar volatilization behavior. The nickel volatility in H\textsubscript{2}O and CO\textsubscript{2} gasification reaches the highest \( R_v \) value 9.0% and 9.5% at 1100 °C. It is apparent that the difference in the nickel volatility between H\textsubscript{2}O and CO\textsubscript{2} gasification is ignorable, so it can be concluded that the volatility of nickel is not dependent on the gasification atmosphere, but rather mainly on the reaction temperature.
3.3. Occurrence Modes of Vanadium in Pet coke and Its Gasification Char

The transformation of different forms of vanadium in the process of CO\textsubscript{2} and H\textsubscript{2}O gasification can be seen from Figure 5a,b. The mass fraction of vanadium bound to exchangeable and carbonates can be barely detected by ICP-AES. The occurrence of vanadium in pet coke, which is similar to that of nickel, is mainly in organic matter (65.78\%) and residual form (28.29\%), with a small amount of Fe-Mn oxides form, which is consistent with Li et al. [18] who reached the conclusion that the organic and residual forms are the predominant vanadium chemical forms while the content of vanadium occurring in exchangeable, carbonates, and Fe-Mn oxides are negligible for raw pet coke.

For the CO\textsubscript{2} gasification, most of the organic-bound vanadium converts to the residual form of vanadium at 800 °C. The content of the organic-bound vanadium sharply reduces from 65.8\% to 6.0\% and the vanadium bound to Fe-Mn oxides decreases from 5.9\% to 2.2\%, resulting in the residual form vanadium reaching a maximum of 90.8\%. As temperature increases to 1000 °C, the residual form of vanadium begins to decompose, mainly converting into vanadium bound to Fe-Mn oxides, while the other forms remains basically unchanged. When the temperature reaches 1100 °C, the Fe-Mn oxides reach the maximum value of 11.0\%, while the organic-bound vanadium is almost completely decomposed. After gasification at 800 °C, V associated with organic matter sharply drops 59.8\%, while the residual form acutely rises 62.5\% with no content of V volatility emerging. At 1100 °C,
V associated with organic matter decreases 64.7% while the residual form acutely rises 53.3% with a little V volatility (1.8%) emerging. It is obvious that the organic-bound vanadium prefers to form residual fraction rather than volatilize. No vanadium bound to exchangeable is found throughout the whole gasification process.

For the H$_2$O gasification, as shown in Figure 5b the organic-bound vanadium decomposes to the residual form at 800 °C. The organic-bound vanadium sharply reduces from 65.8% to 20.8% and the residual form increases from 28.3% to 71.8%. At 900 °C, the residual form of vanadium decreases by 20.0% and changes to the Fe-Mn oxides and carbonates while the organic-bound vanadium remains consistent. As the temperature increases to 1000 °C, both organic-bound and the residual form of vanadium simultaneously reduce and convert into Fe-Mn oxides while the carbonates remains constant. At 1100 °C, the vanadium bound to carbonates converts to the residual form. The organic-bound vanadium shows a downward trend while Fe-Mn oxides display an upward tendency during steam gasification. No vanadium bound to exchangeable is detected.

Comparing the two kinds of atmosphere, the variation of the forms are obviously different. In the CO$_2$ atmosphere, the dominant conversion is the organic-bound vanadium to the residual form. However, the forms in the steam atmosphere are more diverse, by changing from the organic-bound vanadium to other forms of residue, Fe-Mn oxides and carbonates. In both atmospheres, increasing temperature promotes the decomposition of organic-bound vanadium, but the total amount in the H$_2$O is significantly higher than that in the CO$_2$ atmosphere, indicating that the CO$_2$ gasification atmosphere is favorable for the decomposition of organic-bound vanadium. It is worth noting that the vanadium bound to Fe-Mn oxides are gradually increasing in both atmospheres, but the amount in the steam atmosphere is three times higher than that in the CO$_2$ atmosphere, indicating that the steam atmosphere is conducive to the formation of Fe-Mn oxides. The presence of large amounts of Fe-Mn oxides affects the transformation of various forms in the steam gasification process. At the same time, the total amount of the residual form vanadium in the steam atmosphere is significantly lower than that of the CO$_2$ atmosphere, indicating that the H$_2$O atmosphere promotes the conversion of residual form vanadium into other forms.

As the temperature increases, the volatilization characteristic of vanadium gradually increases during H$_2$O and CO$_2$ gasification, as described in Figure 6. It is apparent that H$_2$O and CO$_2$ gasification display similar vanadium volatilization behavior. The vanadium volatility in H$_2$O and CO$_2$ gasification reaches the highest value 1.5% and 1.8% at 1100 °C. It is obvious that the difference in the vanadium volatility between H$_2$O and CO$_2$ gasification is ignorable, so it can be concluded that the volatility of vanadium during gasification is not dependent on the gasification atmosphere, but mainly on the reaction temperature. Li et al. [18] found the volatilization of vanadium is strongly dependent on the vanadium in the organic matter in petcoke.
3.4. Ni Speciation in Different Gasification Atmospheres

As FactSage software platform is based on the Gibbs free energy minimization principle and the system is in a thermodynamic equilibrium state, it is mainly used to analyze the conversion of inorganic substances. Calculation results of CO\textsubscript{2} and H\textsubscript{2}O gasification associated with Ni are shown in Figures 7 and 8.

![Figure 7. Calculation results of CO\textsubscript{2} gasification associated with Ni.](image1)

![Figure 8. Calculation results of H\textsubscript{2}O gasification associated with Ni.](image2)

For the CO\textsubscript{2} gasification, the only product is NiS\textsubscript{2} (s) below 200 °C. For the 300–400 °C temperature range, the only product changes to NiS\textsubscript{3}S\textsubscript{2} (s). Above 400 °C, the main product is NiS\textsubscript{3} (s). Li et al. [18] predicted the morphology of vanadium in the gasification of petcoke. As the temperature increased, Ni reacted with S to form NiS\textsubscript{2}, Ni\textsubscript{5}S\textsubscript{8}, Ni\textsubscript{2}S\textsubscript{5}, and Ni\textsubscript{3}S\textsubscript{2}. The melting points of NiS (s) and Ni\textsubscript{3}S\textsubscript{2} (s) are 645 °C and 797 °C. It can be found that the S content decreases with the increase of Ni, while the melting point and stability of Ni\textsubscript{3}S\textsubscript{2} compounds gradually increase. When the temperature reaches 1300 °C, gaseous Ni (g) and NiS (g) are produced. The thermodynamic equilibrium simulation in [27] showed that Ni\textsubscript{3}S\textsubscript{3} was the stable form of nickel until 1000 °C in a reducing gasification atmosphere (containing H\textsubscript{2}S gas). When the temperature was up to 1000 °C, the main stable nickel form was elemental Ni. For the H\textsubscript{2}O atmosphere, the distribution of Ni is basically the same as in the CO\textsubscript{2} atmosphere, but the formation temperature of Ni\textsubscript{3}S\textsubscript{2} (s) and Ni\textsubscript{3}S\textsubscript{2} (s) are lowered by 100 °C. This indicates that steam is favorable to generate the more stable Ni\textsubscript{3}S\textsubscript{3} compound, thereby suppressing the volatilization of the Ni-containing compound, which is identical with the volatility result of Ni in Figure 4. When the
temperature rises to 1400 °C, there is some solid elemental Ni apart from gaseous Ni (g) and NiS (g). Ni element does not react with other metal elements at this temperature. There are more elemental Ni (s) at high temperature in the H2O atmosphere than that in CO2 atmosphere, which is also consistent with the residual content of Ni at 1100 °C in the experiment (Figure 3).

When comparing the experimental data (Figure 4) with the simulation results, it is found that there is reasonable agreement on the volatilization characteristic of Ni. In the gasification process, nickel is mainly present in the form of Ni3S2 (with a low melting point of 797 °C). When the temperature reaches 800 °C, the volatilization of Ni species occurs, which can also be verified by the Rv changes in Figure 4. Wang et al. [7] investigated the corrosion and fouling in syngas cooler tubes and reported the Ni3S2 would be vaporized instantly during the devolatilization process due to the high temperature. It can be deduced that the main volatilization component of Ni comes from the Ni3S2 in the high temperature range (800–1100 °C).

3.5. V Speciation in Different Gasification Atmospheres

Figures 9 and 10 show thermodynamic calculation results of vanadium under CO2 and H2O gasification process. The results, under the gasification conditions of 800–1100 °C, indicate that the gaseous vanadium compounds formed in the two atmospheres are almost the same, revealing that the atmosphere has little effect on the distribution of V species at different temperatures, which proves that the volatility of V during gasification does not depend on the gasification atmosphere. Below 200 °C, the main product is V3O5, and it changes to V2O3 as the temperature rises from 300 °C to 1500 °C. Below 1300 °C, there is no volatility of V, which is consistent with the experiment results of Rv in Figure 6. Above 1300 °C, a little VO2 gas is generated (less than 5% of the total amount of V). However, V2O3 is still the main product in the gasification process of pet coke. V does not react with other metal elements in this temperature range. The migration and conversion mechanism of V in the process of pet coke gasification is similar to that of coal gasification and the conclusions agree well with some literature [28,29] about the presence of vanadium in coal gasification reaction. Frandsen et al. [28] found that the stable V species was V2O3 (s) and the volatilization of VO2 (g) emerges when the temperature exceeded 1525 °C. Bunt et al. [29] found that V2O3 remained stable while VO2 gas (0.1%) was observed above 1225 °C.

![Figure 9. Calculation results of CO2 gasification associated with V.](image-url)
When comparing the experimental data (Figure 4) with the simulation results, it is found that there is reasonable agreement on the volatilization characteristic of V. References [17,18,30] illustrate that the occurrence forms of V in gasification char were mainly $V_2O_3$, Fe$\text{V}_2O_4$ and other forms of minerals, of which there might be the following reaction during the gasification:

$$2V_2O_3 + CO + Fe_2O_3 \rightarrow 2FeV_2O_4(s) + CO_2$$  \hspace{1cm} (3)

Thermodynamic equilibrium calculation results show that the content of $V_2O_3$ (s) in H$_2$O and CO$_2$ gasification remain stable at 0.004 mol from 300 °C to 1300 °C, but it begins to decrease at 1400 °C in both H$_2$O and CO$_2$ gasification. The modeled results of the content of $V_2O_3$ (s) in H$_2$O gasification at 1400 °C and 1500 °C is 0.00396 mol and 0.00371 mol while that in CO$_2$ gasification is 0.00398 mol and 0.00385 mol. There is an inflection point at 1400 °C which shows the trend of the change of V compound. Therefore, the content of $V_2O_3$ in H$_2$O gasification is lower than that in CO$_2$ gasification at high temperature, mainly because high concentration of CO$_2$ inhibits the positive reaction (3), resulting the increase of residual V and decrease of Fe-Mn oxides, which is also consistent with the residual content of V at 1100 °C in the experiment (Figure 5). Combined with the occurrence patterns of vanadium, the proportion of the residual decreases and the volatile fraction increases as the temperature rises, which is also consistent with the simulation results trend. This shows that the simulation results can predict well the conversion trend of V.

3.6. Fouling, Slagging and Erosion Tendency Assessment for Ni and V Occurrence Modes

Many researchers [31–33] have found that the behavior and effects of heavy metals could not be predicted and explained by their total amount in the environment, but rather were related to the distribution of forms and the differences in migration and transformation behaviors of the various forms, which determined the volatility at high temperatures. Zhang et al. [31] concluded that the exchangeable, acid-soluble and organic matter as soluble form could be leachable under appropriate natural conditions, while insoluble forms, including the Fe-Mn oxides and residue form, were defined as the form that could not be leached. The analysis of morphological changes of heavy metals in the process of fly ash heating indicated that the soluble form of minerals decreased with increasing temperature, and the total of residual and Fe-Mn oxides form increased accordingly, summarizing that high temperature benefits the stability of minerals [31]. Liu et al. [32] found a relatively uniform distribution of five forms of Ni in dry sludge which were mainly distributed in the Fe-Mn oxides form and residue form after thermal treatment. With the increase of temperature, the ratio of state to unstable state increased from 0.4:1 in dry sludge to a maximum of 10.4:1 at 700 °C, indicating that
Ni was present in a more stable form after thermal treatment. Zorpas et al. [33] found that significant conversions of minerals converted to a less mobile form or released by vaporization during heating process. Wang et al. [7] researched the corrosion and fouling in syngas tubes, and found that the insoluble form was relatively stable and it could not be easily volatilized during the heating process, suggesting the problem of being more inclined to the erosion on the metal tube wall. Correspondingly, the soluble form was more likely to cause fouling and slagging.

Therefore, it can be seen from Figures 3 and 5 that during the gasification reaction, V and Ni dominate the erosion problem under the CO\(_2\) atmosphere. The V and Ni insoluble mass fractions are not affected by the temperature change, remaining basically at 93% and 85%, respectively. Compared with the CO\(_2\) atmosphere, the contribution rate of V and Ni to the fouling and slagging is obviously increased under the H\(_2\)O atmosphere, but this impact gradually weakens with the increase of temperature. The influence of atmosphere to the fouling and slagging problems is about the same at the temperature of 1100 °C. In addition, it can be noticed that the erosion problem caused by V is more serious than that by Ni under the two atmospheres, especially in the CO\(_2\) atmosphere, so attention should be paid to the ratio of CO\(_2\) concentration during actual operation.

4. Conclusions

The migration and transformation of V and Ni in high sulfur petcoke during CO\(_2\) and H\(_2\)O gasification were investigated in the range from 800 °C to 1100 °C in a laboratory-scale fixed-bed reactor. The V and Ni volatilization behavior, migration and transformation behavior, and distribution of occurrence modes were studied. The main conclusions were drawn as follows:

1. The organic matter and residual forms were the dominant chemical forms of both V and Ni in pet coke and the Fe-Mn oxides only accounted for a small proportion.
2. The volatilization of V and Ni was not dependent on the gasification atmosphere, but mainly on the reaction temperature.
3. The CO\(_2\) atmosphere was conducive to the conversion of the organic-bound nickel to the residual form Fe-Mn oxides. The H\(_2\)O atmosphere favors the formation of diverse forms.
4. Factsage results confirmed the H\(_2\)O atmosphere was favorable to suppress the volatilization of Ni-containing compounds. More residual content of V was formed in the CO\(_2\) gasification.
5. The V and Ni mainly caused erosion problems under the CO\(_2\) atmosphere while the fouling and slagging obviously increased under the H\(_2\)O atmosphere, with this impact being gradually weakened with the increase of temperature. The erosion problem caused by V was more serious than that caused by Ni, especially in the CO\(_2\) atmosphere.

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