Kinetic and thermodynamic investigations of CO₂ gasification of coal chars prepared via conventional and microwave pyrolysis

Peng Jiang¹ · Yang Meng¹ · Ziyao Lu¹ · Lan Xu¹ · Gang Yang² · Xiang Luo² · Kaiqi Shi² · Tao Wu¹,²

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Abstract This study examined an isothermal CO₂ gasification of four chars prepared via two different methods, i.e., conventional and microwave-assisted pyrolysis, by the approach of thermogravimetric analysis. Physical, chemical, and structural behaviours of chars were examined using ultimate analysis, X-ray diffraction, and scanning electronic microscopy. Kinetic parameters were calculated by applying the shrinking unreacted core (SCM) and random pore (RPM) models. Moreover, char-CO₂ gasification was further simulated by using Aspen Plus to investigate thermodynamic performances in terms of syngas composition and cold gas efficiency (CGE). The microwave-induced char has the largest C/H mass ratio and most ordered carbon structure, but the smallest gasification reactivity. Kinetic analysis indicates that the RPM is better for describing both gasification conversion and reaction rates of the studied chars, and the activation energies and pre-exponential factors varied in the range of 78.45–194.72 kJ/mol and 3.15–102,231.99 s⁻¹, respectively. In addition, a compensation effect was noted during gasification. Finally, the microwave-derived char exhibits better thermodynamic performances than the conventional chars, with the highest CGE and CO molar concentration of 1.30% and 86.18%, respectively. Increasing the pyrolysis temperature, gasification temperature, and CO₂-to-carbon molar ratio improved the CGE.

Keywords Coal char · CO₂ gasification · Microwave pyrolysis · Char properties · Kinetics · Thermodynamic

1 Introduction

Coal is a particularly important energy resource and is currently responsible for approximately 70% of total energy consumption in China (Lan et al. 2018; Zhu et al. 2020). Urgent development of clean coal utilization technology is required because of increasing environmental pollution from coal-fired power plants. Coal gasification, which converts solid coal into syngas at high temperatures, is considered to be the cleanest utilization approach because it offers near-zero sulphur and particulate emissions, high energy efficiency, and flexible chemicals (Attwood et al. 2003; Li et al. 2018). Char reaction with gasification agents is the rate-controlling step of the coal gasification process because it occurs more slowly than water evaporation, pyrolysis, or combustion (Dupont et al. 2011). As a greenhouse gas, CO₂ is the largest contributor to global warming (Fan et al. 2017). Hence, it is critical to implement CO₂ mitigation strategies that can help to alleviate climate change. Fortunately, the employment of CO₂ to gasify coal char for valuable syngas production emerges as a promising approach to reducing CO₂ accumulation. Therefore, studies on the kinetics and thermodynamic performances of char-CO₂ gasification are
essential for the reactor design, control and efficiency (Zhang et al. 2010).

Char-CO$_2$ reactivity is greatly affected by char characteristics, which are determined mainly by coal pyrolysis conditions, of which the pyrolysis temperature is the most important parameter (Wang et al. 2016b). Microwave heating can produce rapid, volumetric, selective, non-contact heating of coal by directly converting electromagnetic energy into thermal energy (Parvez et al. 2019). This is quite different from conventional heating mechanisms in which heat is transferred from the coal surface to the coal interior via conduction (Wu et al. 2015). Previous investigations of microwave pyrolysis of coal were mainly focused on three areas: the effect of microwave pyrolysis on physicochemical properties such as coal grindability and dryness (Ge et al. 2013; Lester et al. 2005; Marland et al. 2000); dielectric properties, interaction mechanisms, and enhancement of coal pyrolysis using microwave absorbers (Liu et al. 2016; Monsef-Mirzai et al. 1995; Peng et al. 2012, 2017); and the properties of pyrolysis products such as gaseous material, tar, and char (Abdelsayed et al. 2018; Reddy et al. 2019; Reddy and Vinu 2016). Researchers also considered microwave reactors and their scale-up (Binner et al. 2014; Salema and Ani 2012). It was demonstrated that microwave pyrolysis showed more gaseous and less tar, high quality liquid fuels and more energy-efficient than conventional pyrolysis (Abdelsayed et al. 2018; Reddy and Vinu 2016).

However, studies on the structure, gasification reactivity, and kinetics of microwave-assisted coal char are rarely reported. Abdelsayed et al. (2018) investigated the effects of pyrolysis temperatures and microwave heating on product distributions and char structure changes of Mississippi coal char and tested the combustion reactivity using a non-isothermal thermogravimetric analysis (TGA). More recently, Liu et al. (2020) analysed the detailed evolutions of char structures and functional groups in low-temperature microwave-prepared char from Zhundong coal. The combustion reactivity of the microwave char was also performed. Results showed that microwave-induced char had a higher burnout temperature than that of conventional pyrolysis char. Nevertheless, little research has been performed on the gasification reactivity and kinetics of CO$_2$ iso-thermal gasification of coal chars prepared via microwave pyrolysis. To ensure the implementation of CO$_2$ gasification, the prediction of thermodynamic performances for char-CO$_2$ gasification is of significant importance (Renganathan et al. 2012). Process simulations of biomass gasification using pure or mixer CO$_2$ have been carried out broadly (Cheng et al. 2016; Guizani et al. 2015; Renganathan et al. 2012; Sadhwani et al. 2017). As far as we acknowledged, no study has been conducted in detailed thermodynamic performances of coal char gasification with pure CO$_2$ as a gasifying medium.

In this study, char-CO$_2$ isothermal gasification kinetic behaviours and thermodynamic performances were investigated using thermogravimetric analysis and Aspen Plus, respectively. One char was derived via microwave pyrolysis, while the other three char samples were produced using conventional pyrolysis at different temperatures for the purpose of comparison. In addition, char chemical compositions, structures, and morphologies were analysed. Moreover, SCM and RPM methods were employed to calculate the kinetic parameters of all chars. Furthermore, thermodynamic performances including syngas composition and CGE of char-CO$_2$ gasification were assessed.

2 Materials and methods

2.1 Materials

Pulverised bituminous Qinghai coal was collected from a domestic power plant in China. The air-dried coal was ground and sieved. To avoid heat- and mass-transfer limitations during heating experiments, only particles smaller than 0.106 mm were collected. Conventional pyrolysis was performed in a horizontal tube furnace, where the coal pyrolysis temperatures were set at 1073, 1173 and 1273 K for 30 min under a pure N$_2$ atmosphere, respectively. The collected chars were stored separately in a dryer and labelled Py1073, Py1173, and Py1273, respectively. The microwave-derived char was prepared in a 2.45 GHz multi-mode microwave-cavity from Nanjing Jiequan Microwave Co., Ltd. Approximately 2.0 g of coal was blended with 20.0 g of silicon carbide (used as a microwave receptor to assist in coal pyrolysis) and pyrolyzed at 1173 K for 30 min. Detailed descriptions of microwave pyrolysis can be found in our previous work (Shi et al. 2017). Char derived from microwave pyrolysis was labelled MW.

Proximate and ultimate analyses of the coal and its derived chars were conducted using a thermos balance analyser (NETZSCH STA 449 PC Luxx, Germany) and a VarioE III Element Analyser (GmbH) according to GB/T 212–2008 and GB/T 476–2008, respectively. The results are presented in Table 1.

The morphology of char samples was examined using a SEM (Zeiss Sigma VP, Germany). Char crystal structures were measured using a powder XRD (Bruker D8 advanced A25) with Cu K$_\alpha$ radiation. The chars were scanned from 10° to 80°. The crystals were characterised quantitatively via their inter-layer spacings ($d_{002}$), stacking heights ($L_c$), and average number of crystallites in a stack ($N_{mean}$). Their expressions are illustrated as (Huo et al. 2014),
the gasification temperature at 25 K/min under a pure N₂ (1173, 1223, and 1273 K) were reached, the N₂ flow was replaced with a CO₂ flow (50 mL/min). When the gasification temperatures were evaluated using the reactivity index, the char mass at time $t$ is calculated as:

$$x = \frac{m_0 - m_i}{m_0 - m_{ash}}$$

where $m_0$ is the char mass at the start of gasification, $m_i$ is the char mass at time $t$, and $m_{ash}$ represents the mass of ash in the char.

The reactivity of different chars at various gasification temperatures were evaluated using the reactivity index, $R_{0.5}$, which is expressed as:

$$R_{0.5} = \frac{0.5}{t_{0.5}}$$

where $t_{0.5}$ represents the time required for char conversion of 50%. A higher reactivity indicates a better gasification performance.

### 2.3 Kinetic model

In general, the gasification rate for a heterogeneous reaction can be described as:

$$r = \frac{dx}{dt} = k(T)f(x)$$

$$k(T) = A \exp \left( -\frac{E_a}{RT} \right)$$

where $A$ is the pre-exponential factor; $E_a$ is the activation energy; and $R$ is the universal gas constant, $R = 8.314$ J/(K mol). Here, $f(x)$ is the gasification mechanism function and $x$ denotes carbon conversion.

In this study, the SCM and RPM were adopted as the mechanism functions to fit the experimental data due to their widely application in simulation of char gasification process (Wang et al. 2016a). The SCM considers that the gasification takes place at the char surface and moves inside. The SCM model is described as follows:

$$r = \frac{dx}{dt} = k_{SCM}(1 - x)^{2/3}$$

The RPM assumes overlapping of pore surfaces. The gasification rate is shown as:

$$r = \frac{dx}{dt} = k_{RPM}(1 - x)\sqrt{1 - \psi \ln(1 - x)}$$

where $\psi$ is a structural parameter that is calculated using a regression method as follows:

$$\frac{t_{50}}{t_{0.8}} = \frac{\sqrt{1 - \psi \ln(1 - x)} - 1}{\sqrt{1 - \psi \ln(1 - 0.8)} - 1}$$

The apparent rate constants of $k_{SCM}$ and $k_{RPM}$ can be obtained from the linear fit of the experimental data with the following expressions of SCM and RPM, respectively.

$$3 \left[ 1 - (1 - x)^{1/3} \right] = k_{SCM}t$$

$$\left( \frac{2}{\psi} \right) \left[ \sqrt{1 - \psi \ln(1 - x)} - 1 \right] = k_{RPM}t$$

After determination of the reaction rate constant, the activation energy and pre-exponential factor are determined by plotting $\ln k$ and $1/T$ via the following equation:

$$\ln k = -\frac{E_a}{RT} + \ln A$$
2.4 Aspen plus simulation

Considering the experimental char-CO<sub>2</sub> gasification temperature varied from 1173 to 1273 K, which are quite suitable for the operating temperature of a typical fluidized-bed gasifier. To evaluate the syngas composition and cold gas efficiency, thermodynamic modelling of char gasification using CO<sub>2</sub> was carried out using Aspen Plus (Li et al. 2014). Detailed simulation descriptions can be found in our previous work (Jiang et al. 2019, 2020). The CGE represents conversion of the char energy content to the lower heating value of syngas, as defined by Eqs. (14) and (15) (Renganathan et al. 2012).

\[
CGE = \frac{LHV_{syg}}{LHV_{char}}
\]  

\[
LHV_{syg} = n_{H_2} LHV_{H_2} + n_{CO} LHV_{CO}
\]

where \( n \) and \( LHV \) refer to the molar flow rate and lower heating value, respectively. The subscripts \( H_2, CO \), and char represent the corresponding species.

3 Results and discussion

3.1 Char characterization

As shown in Table 1, both the volatile and moisture contents of the conventional chars decrease as the pyrolysis temperature increases. In contrast, fixed carbon is observed to have a positive relation with the pyrolysis temperature. As the temperature varies from 1073 to 1273 K, the carbon content and C/H mass ratio increase from 75.2% to 76.5% and from 29.1 to 62.0, respectively. This is expected since high temperatures favour hydrocarbon cracking, which leads to more complete devolatilization. The MW char exhibits the lowest volatile, moisture, and hydrogen contents, but the highest fixed carbon content and C/H ratio. This is mainly determined by the nature of microwaves, which directly convert electromagnetic energy into thermal energy at the centre of the char, thus producing a faster heating rate than conventional heating (Wu et al. 2015). Consequently, more volatile components inside the char are released at high temperatures and higher fixed carbon contents and C/H ratios are obtained.

The crystallinities and structural parameters of the raw coal and pyrolyzed chars were investigated via XRD. The results are displayed in Fig. 1 and Table 2, respectively. Several sharp crystalline diffraction peaks were observed, such as those at 20.8° and 26.6°, which represent the inorganic mineral SiO<sub>2</sub>. In addition, a broad diffraction peak is noted at the 20° angle from 20° to 30°. It corresponds to the (002) carbon crystallite band. Chars prepared at different pyrolysis temperatures exhibit similar diffraction peaks, but the intensity of the (002) diffraction peak increases slightly with the pyrolysis temperature. These phenomena suggest that the microcrystalline structure is prone to becoming ordered. The structural parameters \( d_{002}, L_c, \) and \( N_{mean} \) are employed to quantify the crystal characteristics of all samples as displayed in Table 2. MW char exhibits the smallest \( d_{002} \) of 3.47 Å, which is similar to that of the well-ordered graphite (3.354 Å). Increasing the pyrolysis temperature reduces the \( d_{002} \) of conventional chars from 3.72 Å to 3.65 Å. The \( L_c \) and \( N_{mean} \) increase
from 13.8 Å to 17.43 Å and from 4.71 to 5.77, respectively, with the pyrolysis temperature from 1073 to 1273 K. In addition, the MW char exhibits the highest $L_c$ and $N_{\text{mean}}$, suggesting greater crystallinity. Increasing the pyrolysis temperature can enhance cross-linking via dehydration and decarboxylation and even the dehydrogenation and aromatization reactions, resulting in the increasing ordered structures and also creating new ordered carbons (Abdel-sayed et al. 2018). The microwave-induced char has the most ordered structure due to hot-spot formation under microwave heating. Such hot-spots can result in temperatures that are far higher than those in bulk char (Liu et al. 2019). As a consequence, MW char is more ordered and more thermally stable.

The morphological characteristics of the raw coal and its derived four chars are revealed by the SEM images presented in Fig. 2. The raw coal clearly exhibits a non-porous, blocky shape with a rough surface. The char samples produced via conventional pyrolysis are similar, but more pores and cavities are detected as the pyrolysis temperature increases, as shown in Figs. 2b–d. Besides, the surfaces are prone to be smooth and some microspheres appear as the pyrolysis temperature increases. Due to the release of volatile, some surface pores are formed on the surfaces and the addition of pyrolysis temperature leads particles to fuse and minerals to melt, resulting in the formation of smooth surfaces and microspheres.

In terms of MW char in Fig. 2e, its surface presents a more open structure due to the crack of internal small pores caused by rapid heating expansion. Besides, different size of microspheres with diameter varying from 0.2 to 1 μm are exhibited. This is driven primarily by high localised temperatures in the hot-spots formed via microwave heating (Abdel-sayed et al. 2018). As a consequence, minerals are melted to form such microspheres.

### Table 3 Summary of reactivity index ($R_{0.5}$) values

| Sample  | $R_{0.5}$ (10$^3$/s) |
|---------|----------------------|
|         | 1173 K | 1223 K | 1273 K |
| Py1073  | 1.35   | 1.74   | 2.37   |
| Py1173  | 1.07   | 1.51   | 2.26   |
| Py1273  | 0.68   | 1.48   | 1.95   |
| MW      | 0.39   | 1.08   | 1.92   |
3.2 Kinetic parameters

The reactivity of chars is quantified using the reactivity index, $R_{0.5}$, as detailed in Table 3. As the pyrolysis temperature increases from 1073 to 1273 K at a gasification temperature of 1173 K, $R_{0.5}$ decreases from 1.35 to 0.68. This indicates that the Py1073 char has better gasification reactivity. Similar results can be found at different gasification temperatures. Meanwhile, $R_{0.5}$ has a positive relationship with the gasification temperature. As the conventional char pyrolysis temperature increases from 1173 to 1273 K, the reactivity index increases by 0.75–1.86 times. This suggests that higher gasification temperatures help char gasification. Microwave induced char has a smaller $R_{0.5}$ than conventional chars formed at the same gasification temperature. Increasing the char preparation temperature not only decreases the quantity of volatile matter, but also increases the extent of cross-linking. As a result, the C/H ratio decreases and the carbon structure becomes more ordered, as seen in Table 1 and Fig. 1, respectively. Hence, the reactivity of char gasification is ranked: Py1073 > Py1173 > Py1273 > MW.

The determination of kinetic parameters include reaction constant, pre-exponential factor and activation energy. By plotting $3[1-(1-x)^{1/3}]$ and $(2/\psi)[\sqrt{1-\psi\ln(1-x)}-1]$ versus time ($t$), the reaction constants of $k_{SCM}$ and $k_{RPM}$ are obtained as the slopes of linearised curves at various gasification temperatures. Figure 3 presents a calculating example for the determination of $k_{SCM}$ and $k_{RPM}$ for the Py1073 char. It is worth noting that the value of structural constant $\varphi$ is determined by plotting of $(t/t_{0.8})$ as a function of conversion as shown in Eq. (10) for chars exposed to various gasification temperatures. The $\varphi$ values are regressed to be 8.2, 7.5, 2.8, and 2 for Py1073, Py1173, Py1273, and MW, respectively. This indicates that MW undergoes less pore development during gasification. This result is in accordance with the finding by Liu et al. (2020) that conventional pyrolysis is more conducive to pore development than microwave treatment. Table 4 summarises the rate constants for the two kinetic models. The coefficients of determination ($R^2$) are also listed to show the effectiveness of fitting. As shown, the $R^2$ exceeds 0.99 in all cases suggesting...

![Figure 3](https://via.placeholder.com/150)

**Table 4** $k$ values obtained via linear fits of the experimental data and the coefficients of determination

| Char sample | Model | Reaction rate constant $k$ ($10^{-4}/s$) | 1173 K | $R^2$ | 1223 K | $R^2$ | 1273 K | $R^2$ |
|-------------|-------|----------------------------------------|--------|-------|--------|-------|--------|-------|
| Py1073      | SCM   | 18.9                                   | 25.5   | 0.9967 | 35.3   | 0.9952 |
|             | RPM   | 10.2                                   | 13.9   | 0.9997 | 19.2   | 0.9984 |
| Py1173      | SCM   | 15.1                                   | 22.1   | 0.9941 | 35.9   | 0.9883 |
|             | RPM   | 8.4                                    | 12.3   | 0.9995 | 19.8   | 0.9976 |
| Py1273      | SCM   | 8.2                                    | 19.1   | 0.9992 | 27.6   | 0.9946 |
|             | RPM   | 6.73                                   | 15.1   | 0.9994 | 20.6   | 0.9974 |
| MW          | SCM   | 2.40                                   | 6.14   | 0.9985 | 11.7   | 0.9982 |
|             | RPM   | 2.11                                   | 5.26   | 0.9985 | 10.1   | 0.9985 |
excellent correlation. The resulting values of $k_{\text{SCM}}$ and $k_{\text{RPM}}$ are different for the same char under the same gasification temperature. As the gasification temperature increases, both $k_{\text{SCM}}$ and $k_{\text{RPM}}$ increase by about 1.86–4.88 times. Nevertheless, given a particular gasification temperature, both $k_{\text{SCM}}$ and $k_{\text{RPM}}$ decrease with the pyrolysis temperature, indicating the reduction of reactivity. Based on the calculated $k$ values, the activation energy and pre-exponential factor are determined using the Arrhenius equation shown in Eq. (13). Figure 4 shows plots of ln$k$ versus $1/T$ using various models. Clearly, there is a good linear fit between ln$k$ and $1/T$.

Table 5 summarises the kinetic parameters calculated using the slopes and intercepts in Fig. 4. There are slight differences between the $E_a$ and $A$ values obtained via the SCM and RPM methods. In addition, the pyrolysis temperature and use of microwave heating for char preparation have significant effects on the $E_a$ and $A$, which vary from 78.45 to 194.72 kJ/mol and from 3.15 to 102, 231.99 s$^{-1}$, respectively, using the RPM approach. It is also noticeable that there exists a “compensation effect” as the increase of $A$ when $E_a$ increases (Xu et al. 2019).

To find the best imitative gasification reaction model, the carbon conversion is calculated for all chars as a function of the gasification time at various gasification temperatures. Figure 5 compares the conversions predicted using the SCM and RPM methods to the experimental values. Visually, both models fit the conversion well. However, the RPM prediction is better than the SCM prediction, as the latter exhibits a relatively large discrepancy at times shorter than 800 s, as shown clearly in Fig. 5a, b. To quantify the effectiveness of fitting, both the $R^2$ and root mean square error (RMSE) are calculated for all chars and are displayed in Table 6. The $R^2$ of SCM conversion prediction varies from 0.955 to 0.994, while $R^2$ for RPM prediction ranges from 0.992 to 0.999. In addition, the RMSE values for RPM are between 0.0084 and 0.0251, but the RMSE varies from 0.016 to 0.067 for the SCM. This means that the RPM is the most suitable model for describing gasification conversion.

Variation of $r$ in terms of $x$ is also calculated using both the SCM and the RPM. The predictions are compared to experimental data in Fig. 6. The reaction rate increases until it reaches its maximum at approximately 0.15–0.40. This is followed by a decrease as conversion continues. The reaction rate decrease is attributed primarily to overlapping of inner pores, which leads to reduction of the

| Char sample | SCM $A$ (s$^{-1}$) | RPM $A$ (s$^{-1}$) | SCM $E_a$ (kJ/mol) | RPM $E_a$ (kJ/mol) |
|-------------|-------------------|-------------------|-------------------|-------------------|
| Py1073      | 5.25              | 3.15              | 77.47             | 78.45             |
| Py1173      | 88.49             | 44.25             | 107.28            | 106.23            |
| Py1273      | 4817.45           | 1187.97           | 151.39            | 139.65            |
| MW          | 148,301.11        | 102,231.99        | 197.05            | 194.72            |
reaction surface area and active points (Wang et al. 2016a; Xu et al. 2019). The RPM fits the reaction rate data better than the SCM. Table 6 also shows the $R^2$ and RMSE values used to quantify reaction rate prediction effectiveness. For a given case, the RPM offers a higher $R^2$ and lower RMSE than its counterpart. Hence, the RPM approach is the best for capturing the reaction rate.

### 3.3 Thermodynamic analysis

As previously noted, the influences of the pyrolysis temperature and pyrolysis approach have significant impacts on the char structure and kinetic parameters. To exhibit the char-$\text{CO}_2$ gasification performances including syngas composition and cold gas efficiency under typical fluidized gasifiers using the above four derived chars as the raw materials, Aspen Plus simulation are carried out under the gasification at 1273 K and a $\text{CO}_2$-to-carbon molar ratio ($\text{CO}_2/C$, which is defined as the molar ratio between $\text{CO}_2$ fed to gasifier and carbon content in the char) of 1. The simulation result is depicted in Fig. 7a. The molar concentration of CO exceeds 82% and the molar concentrations of $\text{CO}_2$ and $\text{H}_2$ are approximately 8% and 4%, respectively, in the syngas. When the pyrolysis temperature increases from 1073 to 1273 K, the CO fraction gradually increases from 82.09% to 85.58%, while both the $\text{CO}_2$ and $\text{H}_2$ fractions decrease moderately. The microwave-induced char produces the highest CO concentration of 86.18% and smallest quantities of $\text{CO}_2$ and $\text{H}_2$. This is primarily because the MW sample has the largest C/H ratio, followed by Py1273, Py1173, and Py1073. Given fixed gasification conditions, the higher carbon content in the char generates more CO specie and thus produces a larger CO fraction. The CGE has the similar change tendency as the CO concentration, changing slightly from 1.27 to 1.30. The reason is mainly due to the addition of CO concentration and the LHV of CO is larger than that of $\text{H}_2$. Consequently, an increased CGE is expected according to Eqs. (14) and (15). It is clear that the CGE value exceeds 1 because syngas has a larger LHV than the original char caused by the conversion of $\text{CO}_2$ to CO.

Figure 7b presents the influence of the gasification temperature on gasification performances for the char Py1073 at $\text{CO}_2/C = 1$. Both the CO molar fraction and CGE increase slightly from 88.69% to 88.78% and from 1.266 to 1.267, respectively, when the gasification
temperature increases from 1173 to 1273 K. In terms of H₂ concentration, it exhibits a moderate decreasing trend from 7.21% to 7.12%. The increased gasification temperature helps the endothermic Boudouard reaction (C + CO₂ → CO), which leads to enhancement of char-CO₂ gasification. Besides, the addition of temperature inhibits the exothermic water gas shift reaction (CO + H₂O → H₂ + CO₂). Hence, the CO concentration increases and the H₂ concentration is reduced. Since CO production is favoured, the CGE increases with the gasification temperature.

The CO₂/C ratio is an important parameter because it directly determines the char conversion ratio and syngas composition. Figure 7c shows the effect of changing the CO₂/C ratio from 0.5 to 1.2 at a gasification temperature of 1273 K when testing the char of Py1073. Upon increasing CO₂/C, the CO concentration initially increases and then decreases moderately. Its maximum fraction is 93.8% at CO₂/C = 0.94. Nevertheless, increasing CO₂/C from 0.5 to 1.2 reduces the H₂ concentration from 13.0% to 5.7%. The CGE first increases from 0.74 to 1.22 and then is constant at 1.26 when CO₂/C = 0.94. When CO₂/C < 0.94, the carbon in the char is not fully gasified. The increased CO₂ flow rate contributes to increasing the CO concentration via the Boudouard reaction. Meanwhile, the H₂ concentration decreases because of the reverse WGS reaction. When CO₂/C > 0.94, the molar flowrate of CO remains unchanged. However, continued addition of CO₂ lowers both the CO and H₂ mol fractions after all of the carbon in the char is gasified with CO₂. As a consequence, the CGE first increases and then levels off.

### 4 Conclusions

This study investigated the char structure, morphological evolution, kinetics, and thermodynamics of coal char-CO₂ gasification using XRD, SEM, TGA, and Aspen Plus. Three chars were prepared using conventional heating conditions at 1073, 1173 and 1273 K, while one char was derived via microwave pyrolysis at 1173 K. The main conclusions are as follows.

1. Increasing the pyrolysis temperature enhanced the C/H mass ratio and crystallinity in the char. The microwave-induced char had the highest C/H ratio and most ordered carbon structure. Clear microspheres were observed in the MW char due to hot-spot formation.

2. During gasification, the MW char was less reactive than conventional chars.

3. The kinetic parameters were determined using the SCM and RPM methods. Comparison of R² values indicated that the RPM was better at fitting the gasification conversion and reaction rate experimental data than the SCM. The activation energy and pre-exponential factor were in the range of 78.45–194.72 kJ/mol and 3.15–102,231.99 s⁻¹, respectively. A compensation effect was also noted during the gasification process.

4. The MW char had the best thermodynamic performance, with the highest cold gas efficiency of 1.3 and CO molar concentration of 86.18%. Increasing
the pyrolysis temperature, gasification temperature, and CO₂-to-carbon molar ratio could enhance the cold gas efficiency.

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Conflict of interest The authors declare no conflicts of interest.
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