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1. Introduction

Coal is one of the most commonly used fossil energy resources in the world.1 However, spontaneous combustion of coal is commonplace in many coal-producing countries, resulting in not only safety problems and resource waste2-4 but also ecological damage and environmental pollution.^{5,36} Therefore, understanding the characteristics of coal spontaneous combustion and developing preventative technology is crucial.6 Conventional techniques adopted to prevent and control coal spontaneous combustion are irrigation with water and loess mud.7 However, water and loess mud are unable to completely cover areas affected by the spontaneous combustion of coal. Consequently, a series of coal fire prevention and extinguishment materials, such as cellular grout, three-phase foam, gel foam, and suspension sand have been developed.^{8,37} CO₂ has not only a density greater than air but also an effective flame retardant effect. As a result, pouring CO2 into a goaf will completely bury the lower area, which is beneficial for the prevention and control of coal fires.⁹ Therefore, CO₂ firefighting technology has been developed, and used in the prevention and control of coal fires. Therefore, the inhibiting effect of CO_2 on

Inhibiting effect of CO₂ on the oxidative combustion thermodynamics of coal

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A thermal analysis experiment was conducted in $O_2/N_2/CO_2$ and O_2/N_2 atmospheres (O_2 concentrations were 21, 14, 8, and CO_2 concentrations were 0, 39, 46, 52) to investigate the thermal behavior of coal oxidation and combustion. Results demonstrated that an elevated CO_2 concentration or decreased O_2 concentration had a delaying effect on the thermogravimetric analysis and differential scanning calorimetry (DSC) curves; moreover, the characteristic temperatures were substantially augmented. When the O_2 concentration was 21 vol%, the total heat released by coals A (highly volatile bituminous coal) and B (anthracite coal) decreased by 5.8% and 4.1%, respectively, after CO_2 addition. The comprehensive combustion performance index was also lowered. The DSC curve can be divided into two exothermic peaks, and the ratio of the peak 1 to peak 2 areas decreased with the addition of CO_2 , which indicated that CO_2 inhibited the oxidation of the active functional groups of coal structures. Apparent activation energy in $O_2/CO_2/N_2$ was less than that in O_2/N_2 .

spontaneous combustion of coal under different conditions is the key to improving and using CO_2 firefighting technology. But previous publications have focused on the characteristics of coal oxidation and combustion, and relevant experimental techniques such as thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) have been increasingly used.¹⁰⁻¹² Rotaru et al.¹³ discovered that different types of coal demonstrated different apparent activation energies in the process of spontaneous combustion. Deng et al.14 determined that a large difference in activation energy existed between the oxidation and combustion stages. The study of oxygen-enriched combustion characteristics involves examining the impact of CO2 on coal combustion.15 Liu et al.16 combusted two chars in O_2/CO_2 and O_2/N_2 atmospheres and found that substituting CO2 for inert nitrogen in the oxidizer had a subtle effect on coal combustion. Várhegyi et al.17 demonstrated that during coal combustion in an O2/CO2 atmosphere, the net reaction rate of coal combustion was only proportional to the partial pressure of oxygen. Rathnam et al.18 revealed that combustion reactivity prominently increased at high temperature stages in a $2\% O_2/$ CO_2 atmosphere compared with a 2% O_2/N_2 atmosphere Zhou et al.¹⁹ discovered that CO₂ has a delayed effect on the maximum mass-loss rate and burnout temperature in the coal combustion process. Therefore, the influence of CO2 on characteristics of coal oxidation and combustion should be further study.

To further examine the inhibiting effect of CO_2 on the characteristics of coal oxidation and combustion in different atmospheres, this study used the TGA and DSC methods to analyze the thermal behaviors and variation of apparent activation energies in $O_2/N_2/CO_2$ and O_2/N_2 atmospheres. The main

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purposes of this research were to: (1) investigate the thermal behavior in different atmospheres; (2) analyze the effect of CO_2 and oxygen concentration on apparent activation energies during oxidation and combustion of coal; (3) research the influence of CO2 and oxygen concentration on coal oxidation and combustion. The results are useful for understanding the oxidation and combustion characteristics of coal in O2/N2/CO2 and O₂/N₂ atmospheres and can aid development of CO₂ fire prevention technology.

Experiments and methods 2.

Experimental samples 2.1.

Coal A was highly volatile bituminous coal and was collected from the Juve mining area in Shandong Province, China, whereas coal B was anthracite coal and came from the Furong mining area in Sichuan Province, China. The proximate and ultimate analyses of the two samples are listed in Table 1. The samples had a particle size of 0.067-0.079 mm and were stored in an airtight container.

analyzer, STA449F3 (Netzseh, Geronduct the experimental tests. The , 10, and 15 °C min⁻¹. The experid from 30 to 800 °C for each heating of O_2 , N_2 , and CO_2 , were constantly ¹ flow rate. Table 2 presents the gas ume fractions of O_2 , N_2 , and CO_2 .

rate used in the experiments, single rate heating rate methods were activation energy. In addition, the d must determine the mechanism g the activation energy,20 which may of selecting mechanism functions. rate method can avoid the selection to obtain more accurate values of nstance, Kissinger–Akahira–Sunose Vall–Ozawa (FWO) widely used to the energy, as eqn (1) and (2),

$$\log(\beta_i) = \text{Const} - 1.052 \frac{E_{\alpha}}{RT_{\alpha}}$$
(1)

Table 2 Gases supplied when undergoing thermal analysis experiments

Gas	Volume fraction (vol%)								
	1	2	3	4	5	6			
O_2	21	14	8	21	14	8			
CO_2	0	0	0	39	46	52			
N_2	79	86	92	40	40	40			

$$\log\left(\frac{\beta_i}{T_{\alpha,i}^2}\right) = \text{Const} - \frac{E_{\alpha}}{RT_{\alpha}}$$
(2)

According to the analysis of Starink,²¹ KAS method developed by Starink offers more accurate estimates of E_{α} , as described in egn (3):

$$\log\left(\frac{\beta_i}{(T+273.15)_{\alpha,i}}\right) = \text{Const} - 1.0008 \frac{E_{\alpha}}{R(T+273.15)_{\alpha}} \quad (3)$$

The comprehensive combustion performance index (K) was used to comprehensively evaluate the characteristics of coal spontaneous combustion, and is expressed in eqn (2):22

$$K = \frac{(\mathrm{d}w/\mathrm{d}t)_{\mathrm{max}} \times (\mathrm{d}w/\mathrm{d}t)_{\mathrm{av}}}{T_{\mathrm{ig}}^{2} \times T_{\mathrm{b}}}$$
(4)

Results and discussion 3.

Features on TG-DSC tests 3.1.

According to the characteristic temperatures in Fig. 1, the mass variation of coal was divided into three stages: the lowtemperature pyrolysis stage (stage 1), oxygen-adsorption stage (stage 2), and combustion stage (stage 3). The boundary temperatures for stage 2 were T_1 and T_{ig} , where T_1 is the initial temperature when the coal's mass continuously increases and T_{ig} is the ignition temperature.²⁰ Furthermore, T_2 was defined as the temperature at which the maximum exothermic peak is completed on the DSC curve. Stage 1 was from the initial temperature to T_1 . Stage 3 was from T_{ig} to T_2 .

3.2. Mass variation

Fig. 2 illustrates the TG and differential thermogravimetry (DTG) curves, and the characteristic temperatures are given in Table 3 with a heating rate of 5 °C min⁻¹.

A synchronous thermal a many), was employed to c heating rates were set as 5 mental samples were heater rate. The gases, mixtures of supplied at a 100 mL min mixtures with different volv 2.3. Analysis methods According to the heating r heating rate and multi- concluded to calculate the single heating rate methon functions before calculating cause errors in the process However, the multi-heating of mechanism functions activation energy. For in (KAS) method and Flynn–W calculation of activation respectively. ²¹	2.2. Experimental details
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Coal sample	Proximate ar	Ultimate analysis (%, daf.)							
	Moisture	Ash	Volatile matter	Fixed carbon	С	Н	0	Ν	S
Coal A	2.3	9.7	34.85	53.15	85.71	4.58	7.48	1.54	0.69
Coal B	1.4	26.02	8.85	63.73	89.69	3.61	1.71	2.85	2.14



Fig. 1 TG and DSC curves of a coal sample showing the temperature boundaries for different stages. This individual diagram is obtained from the thermal analysis of coal A conversion under 21 vol% $O_2/79$ vol% N_2 and heating rate of 5 °C min⁻¹.

Both the increase of the CO₂ concentration and decrease of the O₂ concentration made the TG/DTG curve move to a high temperature range, as illustrated in Fig. 2. T_{ig} and T_{max} decreased with increasing O2 concentration. For example, when the O_2 concentration was reduced from 21 to 8 vol%, the T_{ig} of coal A increased from 308.4 to 323.1 °C, and the T_{ig} of coal B increased from 309.2 to 325.5 °C (Table 3). The lower the oxygen concentration in the environment, the lower the number of activated oxygen molecules. Therefore, the opportunities of collision between activated oxygen molecules and active functional groups of coal is significantly reduced, and the reaction between coal and oxygen becomes difficult. This finding is in agreement with the reports by Deng et al.,14 Wang et al.,23 and Liu et al.24 When the O2 concentration was constant, increasing the CO₂ concentration had a delay effect on the TG and DTG curves, whereas the characteristic temperatures increased along with CO2 concentration. As Table 3 shows, when the O2 concentration was 21 vol% and the CO₂ concentration was increased from 0 to 52 vol%, the ignition temperature of coal A increased from 323.1 to 325.5 °C and the ignition temperature of coal B increased from 382.3 to 387.4 °C. The diffusion rate of O_2 in CO_2 is lower than that in N_2 , which has an adverse effect on the transportation of O₂ from the gas mixture to the coal particle surface.^{23,25} In addition, CO₂ is much easier for coal to

physically adsorb than O_2 and N_2 .^{26,27} Therefore, adsorption of CO_2 occupies some of the active sites on the coal surface, inhibiting the ability of O_2 and coal to interact.

3.3. Exothermic properties

3.3.1. Differential scanning calorimetry. The DSC and firstorder differential of DSC (DDSC) curves for the two coal samples from 30 to 800 °C were illustrated in Fig. 3. Total heat release is presented in Table 4.

Fig. 3 shows that exothermic peaks occur when the temperature exceeds 200 °C. The exothermic process finished when the temperature was more than 600 °C. The DSC curve of coal exhibited typical bimodal exothermic behavior. However, the first exothermic peak of the DSC curves for coal B was not clear. This phenomenon is caused by differences in the types and quantities of active structures contained in coal. Some active structures, such as alkyl chains, methyl groups, carboxyl groups, and hydroxyl groups, are easily oxidized at lower temperatures.^{28,29} However, the combustion or decomposition temperature of some stable structures, such as aromatic structures, is higher.³⁰ When the concentration of CO₂ was zero, the maximum heat intensity and total heat increased significantly with an increase in the O_2 concentration. When the O_2 concentration was increased from 8 to 21 vol%, the total heat released by coal rose from 17.296 to 19.247 kJ g^{-1} , as listed in Table 4. When the O_2 was sufficient, the coal-oxygen reaction was more intense. With the increased O2 concentration, the coal and O2 were able to have more contact; therefore, the oxidation/ combustion reactions of the coal produced a high proportion of stable oxidation products, increasing heat intensity. When the O2 concentration was kept constant, the DSC and DDSC curves moved to the high temperature region, whereas total heat release decreased substantially with increasing CO2 concentration. As Lee³¹ noted, competitive adsorption occurs in coal when various gases exist. The adsorption of CO2 causes a reduction in the amount of O_2 contacting coal. In addition, more unstable products are formed, leading to an apparent reduction in heat intensity.

3.3.2. Stage characteristics of heat release. The DSC curves for the two samples at a heating rate of 5 °C min⁻¹ and in O_2 : N_2 : $CO_2 = 21$: 79: 0 and O_2 : N_2 : $CO_2 = 21$: 40: 39 atmospheres are illustrated in Fig. 4. The sub-peaks of the DSC



Fig. 2 TG and DTG curves of the two coal samples with the heating rate of 5 $^{\circ}$ C min⁻¹ under different atmospheres, (a) coal A and (b) coal B.

Table 3 Characteristic temperatures during combustion of coal with heating rate of 5 °C min⁻¹

Cample Atmosphere	Coal A			Coal B				
Sample Atmosphere $(O_2 : N_2 : CO_2)$	T_1 (°C)	T_{ig} (°C)	$T_{\max} \left(^{\circ} \mathbf{C}\right)$	T_2 (°C)	T_1 (°C)	$T_{\mathrm{ig}} (^{\circ}\mathrm{C})$	$T_{\max} \left(^{\circ} \mathbf{C}\right)$	$T_2 (^{\circ}C)$
21:79:0	81.3	308.4	482.9	579.5	83.35	367.6	523.2	617.9
14:86:0	79.1	313.6	500.1	591.1	81.45	370.1	539.7	627
8:92:0	83.1	323.1	516.2	616.4	70.25	382.3	567.1	649.2
21:40:39	71.9	309.2	485.4	583.4	84.55	371.7	524.9	618.8
14:40:46	80.6	317.9	499.7	598.7	81.45	378.6	546	634.7
8:40:52	75.7	325.5	523.9	627.5	80.55	387.4	576.5	660.2

curves were fitted using eqn (5) (Gaussian equation). The fitting parameters of the sub-peaks are given in Table 4. Fig. 5 illustrates the ratio of the peak 1 to peak 2 areas.

$$v_{\rm m} = v_0 + \frac{S\left(\left(-4\ln(2)(t-t_{\rm c})^2\right)/w^2\right)}{w\sqrt{\pi/(4\ln(2))}}$$
(5)

The temperature corresponding to the maximum of peak 1 in coal A was in the range of 320-360 °C, whereas it was in the range of 410-440 °C for coal B, as listed in Table 4. Furthermore, the temperature corresponding to the maximum of peak 1 was close to the ignition temperature of coal. Some of the active structures in coal, such as alkyl chains, methyl groups, carboxyl groups, and hydroxyl groups, are oxidized at low temperatures, resulting in a clear oxidative-exothermic peak before coal combustion.³⁰ Peak 2 was caused by the breaking of numerous stable structures (such as aromatic rings) into small molecules that participated in the combustion reaction.³² As illustrated in Fig. 5, the ratio for coal A was clearly larger than that for coal B, which indicates that attribution of peak 1 to the total heat release was greater than coal B. The primary reason for this phenomenon is that the metamorphic grade of coal A was lower than that of coal B, and the molecules of coal A contained numerous alkyl chains, methyl groups, carboxyl groups, hydroxyl groups, and other active structures.33 The ratio decreased with a decrease of the O2 concentration or increase of the CO₂ concentration, which indicates that a low O₂ concentration or high CO₂ concentration inhibits oxidation of some reactive groups in coal.

3.4. Comprehensive combustion performance index

The dependence of K on the O₂ concentration for the two samples is illustrated in Fig. 6. The comprehensive combustion performance index of the two samples decreased with a decrease in the O₂ concentration; furthermore, the higher the O₂ concentration was, the more O₂ participated in coal combustion, leading to the combustibility being strengthened. For the same O₂ concentration, the addition of CO₂ resulted in a significant decrease in the comprehensive combustion performance index. The physical adsorption of CO₂ onto the coal surface occupied active sites, which hindering the physical adsorption and chemical adsorption of O₂ by coal, and further inhibiting coal oxidation and combustion.

3.5. Apparent activation energy

The calculation of the apparent activation energy uses the Starink method, which is a multiple heating rates calculation method. Therefore, a set of data points of multiple heating rates (5, 10 and 15 °C) calculated at intervals of 0.5 from 0.5 to 0.95 was obtained according to eqn (3), and $\ln(\beta/(T + 273.15)^{1.92})$ and 1/(T + 273.15) were calculated at different conversion rates as shown in Fig. 7. The apparent activation energies during stages 2 and 3 for the two coal samples are given in Fig. 8 and 9.

3.5.1. Oxygen-adsorption stage. The physical and chemical adsorption of O_2 and CO_2 is easier and the energy required for the chemical reaction between O_2 and coal is lower at the beginning of the oxygen-adsorption stage. The reaction between coal and oxygen is a step-by-step reactive process in which functional groups are gradually activated and oxidized. At low temperatures, only some active structures are activated and



Fig. 3 DSC and DDSC curves of the two coal samples with the heating rate of 5 °C min⁻¹ under different atmospheres, (a) coal A and (b) coal B.

Table 4 Factors used to fit sub-peaks for DSC curves and total heat release and the maximum of exothermic peak with heating rate of 5 °C min⁻¹

Coal sample	Atmosphere $(O_2 : N_2 : CO_2)$	Peak	ν_0	S	t _c	w	R^2	Q (kJ g ⁻¹)
Coal A	21:79:0	1	0.306	-172.495	66.473	35.291	0.9896	19.247
		2		-162.325	90.641	13.913		
	14:86:0	1	0.156	-158.973	68.084	35.237	0.9939	17.891
		2		-159.263	93.457	15.907		
	8:92:0	1	0.042	-139.887	69.772	34.700	0.9972	17.296
		2		-175.342	97.945	19.815		
	21:40:39	1	-0.141	-162.444	66.061	34.582	0.9889	18.177
		2		-158.456	90.895	14.639		
	14:40:46	1	-0.274	-118.875	67.107	33.344	0.9942	17.893
		2		-150.177	94.420	18.585		
	8:40:52	1	-0.093	-131.845	70.106	34.198	0.9938	17.794
		2		-172.622	99.379	20.963		
Coal B	21:79:0	1	0.038	-103.340	82.755	38.798	0.9972	15.308
		2		-138.154	97.773	12.381		
	14:86:0	1	-0.165	-89.947	83.914	39.612	0.9984	13.812
		2		-141.981	100.100	13.747		
	8:92:0	1	0.004	-68.126	84.428	39.436	0.9967	13.018
		2		-152.792	106.658	16.9603		
	21:40:39	1	-0.200	-106.997	83.199	39.943	0.9976	14.779
		2		-144.890	98.404	12.736		
	14:40:46	1	0.021	-76.145	83.285	37.229	0.9983	13.197
		2		-144.658	102.357	14.770		
	8:40:52	1	-0.081	-68.306	87.135	42.810	0.9948	12.476
		2		-143.390	108.245	17.740		

react with O_2 . Moreover, these oxidation reactions require less reaction energy, resulting in lower apparent activation energy of coal. As the temperature increases, more stable structures are

gradually activated and react with O_2 . These oxidation reactions require more energy, leading to an increase in apparent activation energy. Therefore, we observed that with an increase of



Fig. 4 DSC curves from the pyrolysis domain stage (stage 1) through the combustion domain stage (stage 3) for the two coal samples, (a) coal A in atmosphere $O_2 : N_2 = 21 : 79$, (b) coal A in atmosphere $O_2 : N_2 : CO_2 = 21 : 40 : 39$, (c) coal B in atmosphere $O_2 : N_2 = 21 : 79$, and (d) coal B in atmosphere $O_2 : N_2 : CO_2 = 21 : 40 : 39$, (c) coal B in atmosphere $O_2 : N_2 = 21 : 79$, and (d) coal B in atmosphere $O_2 : N_2 : CO_2 = 21 : 40 : 39$, (c) coal B in atmosphere $O_2 : N_2 = 21 : 79$, and (d) coal B in atmosphere $O_2 : N_2 : CO_2 = 21 : 40 : 39$.

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Fig. 5 Ratio of the peak 1/peak 2 areas



Fig. 6 Relationship between the comprehensive combustion performance index and O_2 concentration.

the conversion rate, apparent activation energy increased. Furthermore, the tendency and value of apparent activation energy changed relative to the conversion rate at all percentages of O_2 in O_2/N_2 or $O_2/CO_2/N_2$, while tendency of apparent activation energy was almost the same, as illustrated in Fig. 8. When the CO_2 concentration was 0, the apparent activation energy decreased with an increase in the O2 concentration. In light of the criteria of competitive reactions, the primary reason for this behavior is that the active structures consume a considerable amount of O_2 . The decrease of the O_2 concentration inhibits the reaction of the stable structures in coal with O₂, which requires more energy. In addition, coal B was anthracite, which provides more surface area for strongly competitive chemisorption,34 resulting in complex changes of the apparent activation energies. When the O_2 concentration was the same, the apparent activation energy of coal oxidation in O₂/N₂ was greater than that in O₂/CO₂/N₂. Liu et al.²⁴ considered chemisorption in O₂/CO₂ conditions to be mainly controlled by O₂ chemisorption at the oxygen-adsorption stage. Moreover, CO_2 competes with O_2 when they are adsorbed by active sites on the coal surface, which hinders the chemical reaction of coal and O_2 . In summary, O_2 competed with CO_2 for active sites; thus, more energy was required.

3.5.2. Combustion stage. The apparent activation energy of coal A first increased and then decreased with an increase in the conversion rate. Coal B exhibited a similar tendency when the O₂ concentration was more than 8 vol%. When the O₂ concentration was 8 vol%, the apparent activation energy of coal B decreased with an increase in the conversion rate, as illustrated in Fig. 9. Temperatures when the apparent activation energy reached the maximum were close to the devolatilization temperature of coal. When the temperature exceeded the devolatilization temperature, a large volume of volatiles evaporated out, leading to an increase in the diffusion resistance of O₂ to the coal pores.³⁵ Then, the apparent activation energy of coal combustion tended to decrease due to the large number of active structures resulting from pyrolysis. Values of the apparent activation energy in N2/O2 atmospheres were larger than those in O₂/N₂/CO₂ atmospheres. The cause of this phenomenon is that competition chemisorption of O₂ and CO₂ on the coal surface during combustion in the O2/N2/CO2 atmosphere, which causes the chemical adsorption process of oxygen in the O₂/N₂/CO₂ atmosphere significantly longer than in the O₂/N₂ atmosphere.²⁴ Consequently, the probability of oxygen contacting with the active structure is reduced.



Fig. 7 Linear relationship between $\ln(\beta/(T + 273.15)^{1.92})$ and 1/(T + 273.15) of coal A in atmosphere $O_2 : N_2 : CO_2 = 21 : 40 : 39$, (a) in oxygenadsorption stage and (b) combustion domain stage.



Relationship between apparent activation energy and conversion during the oxygen-adsorption stage (stage 2), (a) coal A and (b) coal B. Fig. 8



Fig. 9 Relationship between apparent activation energy and conversion during the combustion domain stage (stage 3), (a) coal A and (b) coal B.

Therefore, the combustion rate of coal in the O₂/N₂/CO₂ atmosphere is lower than that in the O_2/N_2 atmosphere (Fig. 2).

4. Conclusions

• With an increase in the CO₂ concentration or decrease in the O2 concentration, the thermal analysis curves of coal were delayed and the characteristic temperatures increased.

• The DSC curve can be divided into two exothermic peaks, and the ratio of the peak 1 to peak 2 areas decreased with the addition of CO2, which indicated that CO2 inhibited the oxidation of active functional groups of coal structures. Furthermore, the total heat release decreased. Higher concentrations of CO₂ or lower concentrations of O₂ resulted in a lower comprehensive combustion performance index.

• At the oxygen-adsorption stage, the apparent activation energy increased with an increase in the conversion rate, whereas it first increased and then decreased at the combustion stage. The apparent activation energy in the $O_2/CO_2/N_2$ atmosphere was less than that in the O_2/N_2 atmosphere.

Nomenclature

- E_{α} Apparent activation energy $(J \text{ mol}^{-1})$
- Ι Constant
- Κ Comprehensive combustion performance index $(\%^2 \min^{-2} \circ C^{-3}))$

Q	Total heat release (kJ mg ⁻¹)
R	Gas constant (J mol ^{-1} K ^{-1}))
S	Peak area (mW min mg^{-1})
Т	Temperature (°C)
T_1	Initial temperature when the coal's mass
	continuously increases (°C)
T_2	Temperature is exothermic completed on DSC curve
	(°C)
$T_{\rm b}$	Burnout temperature (°C)
t _c	Center of the peak time (min)
T_{ig}	Ignition temperature (°C)
ν_0	Baseline of peak (mW mg ⁻¹)
w	Full width at half maximum (min)
$v_{\rm m}$	Change rate of exothermic (min)
$(dw/dt)_{av}$	Average rate of mass loss (% min^{-1})
(dw/	Maximum rate of mass loss (% min^{-1})
dt) _{max}	
α	Conversion rate (%)
в	Heating rate (°C min ⁻¹)

Conflicts of interest

There are no conflicts to declare.

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