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Thermogravimetric Study of the Kinetics of the Reaction C + CO₂ under Pore-Diffusion Control

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Abstract: This study presents experimental studies of charcoal gasification with CO₂ at different heating rates (1, 5, 10, 20, and 50 K min⁻¹). The kinetics of the reaction C + CO₂ under pore-diffusion control is studied. We propose a new method for the proper determination of activation energy during the processing of thermogravimetric curves of porous carbon gasification under conditions of pore-diffusion resistance. The results of the inverse kinetic problem solution are compared with different hypotheses about the regime of the investigated heterogeneous reaction process (kinetic, diffusion, pore-diffusion). The change of reaction regimes from kinetic to diffusion is detected during charcoal gasification at different heating rates. At heating rates of 5–20 K min⁻¹, the values of activation energy of carbon gasification reaction in the carbon dioxide atmosphere, obtained by the proposed method, closely match the data found in the previous studies. The use of diffusion models in the processing of thermogravimetric curves determines the conditions under which conventional kinetic models fail to provide adequate information about the temperature dependence of the heterogeneous reaction rate.

Keywords: gasification; thermogravimetry; thermokinetic analysis; diffusion processes

1. Introduction

According to energy development projections (e.g., Reference [1]), solid fuel (coal and biomass) will remain one of the main energy sources for a long time to come. One of the ways to increase the environmental efficiency of solid fuel use is its gasification. The possibility of combining energy and chemical production (first of all, synthesis liquid hydrocarbons and oxygenates, as well as hydrogen) based on a single thermochemical process opens the prospects for the development of environment-friendly multipurpose plants with chemical energy storage and export [2,3]. The development of gasification technologies requires reliable information on the kinetics of reactions of carbon fuels with gaseous reagents (O₂, CO₂, H₂O).

The observed carbon gasification rate factors often show high variability (e.g., see References [4,5]). It is due to a large number of simultaneously occurring physical and chemical processes that are not considered within the framework of the simplified analysis of processes of thermochemical conversion of fuel in real furnaces and gasifiers. Even for the same chemical reaction under different conditions (e.g., at different heating rates or gas flow rates), different kinetic patterns may be observed. As a rule, it is connected with carbon origin (for example, conditions of raw material pyrolysis) or with the effect of transfer processes.

Gasification kinetics is related to chemical reactions between bound and free forms of carbon and oxygen on the reactive surface [6,7], the state of which is determined by many factors. The type of original material is of key importance [8,9]. However, differences in reactivity of carbon are observed even when it is obtained from the same solid fuel, but under different conditions [10–12]. During pyrolysis, there is particle
cracking under the pressure of released gases, contraction, or, on the contrary, swelling of particles, and bubbling of released gases in the semi-molten state. Higher temperatures facilitate cross-linking between carbon microcrystalline structures, which leads to a decrease in the reactivity of carbon [13,14]. In kinetic modeling of fuel processing, most often carbon gasification reactions are treated as single-stage heterogeneous transformations with an effective kinetic constant that covers the whole set of elementary stages without taking into account the peculiarities of the limiting stages [15].

Carbon dioxide is usually a “secondary” gasification agent that is formed when fuel carbon is oxidized with oxygen. On the other hand, CO₂ is a greenhouse gas that is generated in energy production and chemical technology processes. Therefore, to balance its formation, it is necessary to develop suitable methods for its utilization. In this case, CO₂ can act as a primary gasification agent (for example, in allothermic processes [16]).

There is a wide variety of laboratory reactors for carbon gasification kinetics studies. First of all, it is the drop tube reactor [17,18], and the quartz furnace [19,20]. The fluidized bed reactor makes it possible to achieve a high degree of isothermicity [21]. Fixed bed reactors are used less often because they are most exposed to the weakly controlled effects of transfer processes [22,23]. The least exposed to such influence are thermogravimetric (TG) furnaces whose design, due to high sensitivity, studies thermochemical transformations in small weighed quantities at high rates of heat and mass transfer [24].

An important issue in studying the kinetics of gasification processes is the precise control of temperature fields in the sample. The sample has to be of low thermal inertia, while maintaining, however, its representativeness. As is known from the work on the theory of thermal explosion [25,26], thermogravimetric measurements of exothermic processes require special conditions to avoid overheating, for example, very low rates of heating and dilution with inert filler. Estimates show that even in specialized laboratory devices used to study the kinetics of coal oxidation, it is possible to underestimate the temperature of the sample, due to internal heat generation: In Reference [27], the estimate of the peak temperature difference between the sample and the gas was several tens of degrees; in Reference [28], overheating was more than two hundred degrees at the maximum rate of mass loss. In the study of endothermic processes (such as evaporation and decomposition), deviations in the temperature of the sample from the ambient temperature are possible as well [29]. For more detailed analysis, detailed thermohydrodynamic models of such processes, including chemical heat sources, are being developed [30,31].

When reacting in the diffusion regime, the carbon particle interacts with the reacting gases only by its outer surface, the area of which can be estimated in a rather straightforward way. This proves way more difficult with the internal reacting surface. For example, the processes of pulverized carbon gasification in the atmosphere of carbon dioxide and water vapor even at high temperatures may proceed in pore-diffusion regimes [32,33]. Combustion and gasification at low temperatures (smoldering combustion) also take place in these regions.

Depending on the rate of change of the inner surface, carbon particles behave differently during the reaction: They can burn on the outside, maintaining their density, or, conversely, preserve their volume by reducing their density [34,35]. Oxidation is also affected by the mineral impurities: If the loose ash formed on the surface can be carried away by gas, the dense ash shell creates additional diffusion resistance. The rate of a heterogeneous chemical reaction is extremely sensitive to mass transfer conditions. Even under thermal analysis conditions, when the mass of the weighed quantity is about 10–20 mg, diffusion reaction regimes can be observed [36,37]. The papers [38–40] provide recommendations on the choice of experimental parameters (particle size, mass of samples, gas consumption, heating rate) to prevent transfer effects. In Reference [41], the method of estimation of diffusion influence on sample conversion by carrying out preliminary experiments with naphthalene evaporation to measure mass transfer coefficient was proposed. For thermogravimetric measurements, small particle size is
usually used. In Reference [42], a comprehensive study of the effect of crucible geometry on the pyrolysis and gasification of coals was carried out: The authors detected a significant influence of crucible height on conversion kinetics at high temperatures.

Ya. B. Zeldovich [43] demonstrated that there can be an intermediate pore-diffusion regime between the kinetic and diffusion regimes of the heterogeneous reaction in porous material under certain conditions (high reactivity, high specific surface, large particles). One of the characteristic features of this regime is a twofold reduction of the observed activation energy. Pore diffusion is a typical mass transfer process for carbonous samples [44]. In Reference [45], it is shown that depending on the size of the particles, there are regimes of gasification with different degrees of utilization of the inner surface. Diffusion regimes of conversion under thermal analysis conditions were reproduced with detailed thermohydrodynamic models [46,47]. In Reference [48], a two-dimensional model of heat and mass transfer in a porous carbon sample during gasification under conditions of rapid heating and subsequent maintaining of the constant reaction temperature is proposed. In the present study, the case of a constant heating rate during the whole experiment will be considered.

A decrease in observed (effective) activation energy can be one of the signs of deviation of gasification regimes from occurring in the kinetic region (here, the kinetic region is understood as isothermal conditions with ideal mixing of components). Based on published research data, it is possible to set permissible limits for the activation energy of gasification reactions. Then, for example, it is possible to exclude from consideration obviously unphysical values (too high or too low), which are often given in the studies on the thermogravimetric analysis of solid fuels [49–51].

In the present paper, based on a simple theoretical model, we propose a new method for determining the activation energy of heterogeneous reactions from experimental data with diffusion-induced perturbations. The application of this method is demonstrated by the example of the well-known gasification reaction of porous carbon in carbon dioxide. A technique for the numerical processing of the TG signal is proposed. Results are compared with widely used kinetic models of a one-stage reaction. It is shown that the pore-diffusion model can be used to explain fluctuations in the experimentally observed values of activation energy.

2. Materials and Methods

2.1. Materials

In this study, we used charcoal produced from pine (Bétula). In order to produce the charcoal, we used a muffle furnace, placing an alumina crucible with a lid at 500 °C and holding it for 7 min. Then the crucible was pulled out and cooled down in the desiccator to room temperature. The isothermal holding time is sufficient for the complete devolatilization of the wood biomass. Technical characteristics and elemental composition of pine and charcoal are given in Table 1. The specifications were determined using the relevant standards (ASTM standard E871 (Standard Test Method for Moisture Analysis of Particulate Wood Fuels), E872 (Standard Test Method for Volatile Matter in the Analysis of Particulate Wood Fuels), and D1102-84 (2013) (Standard Test Method for Ash in Wood). The elemental composition was determined using the elemental analyzer (Flash EA 1112, ThermoFinigan, Italy).
Table 1. Technical specifications of raw wood and charcoal.

| Technical Characteristics (% wt., Dry Basis) | Pine      | Charcoal |
|---------------------------------------------|-----------|----------|
| Fixed carbon                               | 10.23     | 58.72    |
| Ash content                                | 0.17      | 1.99     |
| Volatile yield                             | 89.6      | 39.29    |

| Elemental composition (% wt., dry ash-free basis) | Pine | Charcoal |
|--------------------------------------------------|------|----------|
| C                                                | 46.60| 79.84    |
| H                                                | 6.32 | 5.79     |
| O                                                | 47.08| 14.37    |
| Brutto-formula                                   | CH1.62O0.76 | CH0.46O0.14 |

2.2. Thermal Analysis

Experimental studies were conducted with the NETZSCH STA 449 C simultaneous thermal analyzer (Netzsch, Ahlden, Germany). The samples we studied were heated at different rates of 1, 5, 10, 20, and 50 K min⁻¹ in a mixture of argon (purity is 99.9998%) and carbon dioxide (purity is 99.99%). Argon serves as a shielding gas (10 mL min⁻¹). The flow of carbon dioxide was 30 mL min⁻¹. The process of charcoal conversion was studied in a broad temperature range from room temperature to 1273 K. The weight of the samples in all experiments was 9–10 mg. The studies were conducted using a TG holder and alumina crucibles shaped like a flat plate. The sample is placed in a flat crucible, after which, heating and gas flow are turned on (Figure 1). The temperature is automatically regulated by heating the walls. Throughout the experiment, the sample weight and crucible temperature are recorded. After reaching a temperature of 1273 K, experiments were stopped. This is due to limitations on the properties of the furnace materials and the onset of phase transitions in the mineral part of the samples.

A constant heating rate simulates the initial stage of allothermal gasification. Selected heating rates can be observed in low-capacity reactors (for example, in moving bed gasifiers, where the residence time of particles may be up to tens of minutes [52]). The heating rate determines the ratio of the characteristic times of the chemical reaction ($t_R$) and heating ($t_H$). If in the course of heating the dependence of the reaction rate on temperature changes (for example, the transition from the kinetic regime to the diffusion regimes occurs), then at low $t_R/t_H$ ratios, the heterogeneous reaction will proceed closer to the kinetic region, and at large $t_R/t_H$, the reaction will proceed closer to the diffusion region.
This dependence is schematically presented in Figure 2—the higher the heating rate, the smaller the fraction of the sample is converted before transition temperature is achieved.

Figure 2. Schematic dependence of the behavior of TG-curves on the heating rate.

Charcoal has a developed surface, the state of which largely determines the reactivity. It is known that under heating conditions, the reactivity of carbon materials can decrease, due to thermal annealing. However, the studies on this effect as applied to lignocellulosic-origin chars show that it manifests itself at higher temperatures compared to those used in this work [53].

2.3. Kinetic Models for Processing TG Curves

The rate of charcoal gasification is proportional to the area of the reactive surface and the concentration of the gasification agent. The reaction rate constant can usually be represented in the Arrhenius-type form:

\[
\frac{dm_c}{dt} = -k_0 \exp\left(-\frac{E_a}{RT}\right) S_r C_{CO_2}
\]

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\]  

(1)

Here, \(m_c\)—mass of carbon in the sample, \(S_r\)—area of reactive surface, \(m_0\); \(C_{CO_2}\)—concentration of carbon dioxide in gas flow. Equation (1) is often used in modeling solid fuel gasification processes [54]. The dependence of the reaction area on the sample mass is determined by the geometric parameters of the particles; in general, it is possible to assume the formula \(S_r/S_0 = (m_c/m_0)^n\). The linearization of Equation (1) in view of this formula yields:

\[
\ln \left(\frac{dm_c}{dt} \frac{1}{C_{CO_2} m_c^n}\right) = \ln \left(\frac{k_0 S_0}{m_0^n}\right) - \frac{E_a}{RT}
\]

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\]  

(2)

Under the external diffusion regime, the rate of gasification is determined by the rate of oxidizer diffusion to the carbon surface:

\[
\frac{dm_c}{dt} = -\frac{Nu_c D}{L} S_r C_{CO_2}
\]

\[
\frac{dm_c}{dt} = -\frac{Nu_c D}{L} S_r C_{CO_2}
\]  

(3)
Here $N_{uD}$—mass transfer Nusselt number (Sherwood number), $D$—diffusion coefficient, $L$—characteristic crucible size, $S_c$—crucible crucible cross-section. The diffusion coefficient depends on the temperature as its power function: $D = D_0 T_0^p$, where the exponent $p$ is in the range of 1 to 2 [55]. Then linearization is carried out in logarithmic coordinates, not in Arrhenius coordinates:

$$\ln \left( -\frac{dm_c}{dt} \frac{1}{C_{CO_2}} \right) = \ln \left( \frac{N_{uD} D_0 S_c}{L} \right) + p \ln T$$

(4)

$$\ln \left( -\frac{dm_c}{dt} \frac{1}{C_{CO_2}} \right) = \ln \left( \frac{N_{uD} D_0 S_c}{L} \right) + p \ln T$$

Under pore-diffusion regimes, gasification does not occur in the entire porous volume of the sample, but in the diffusion layer, the width of which depends on the ratio of the reaction rate constant and diffusion coefficient (Thiele number [56]). The quasi-stationary rate of the first-order reaction in a layer of particles is given as follows [43]:

$$\frac{dm_c}{dt} = -S_c C_{CO_2} \sqrt{k_0 \exp \left( -\frac{E_a}{RT} \right) DS_p} = -S_c C_{CO_2} \exp \left( -\frac{E_a}{2RT} \right) \sqrt{k_0 DS_p}$$

(5)

$$\frac{dm_c}{dt} = -S_c C_{CO_2} \sqrt{k_0 \exp \left( -\frac{E_a}{RT} \right) DS_p} = -S_c C_{CO_2} \exp \left( -\frac{E_a}{2RT} \right) \sqrt{k_0 DS_p}$$

Here $S_p$—specific surface of particles, m⁻¹. Converting this equation to Arrhenius coordinates produces the following expression:

$$\ln \left( \frac{dm_c}{dt} \frac{1}{C_{CO_2}} \right)^2 = \ln \left( S_p^2 k_0 DS_p \right) - \frac{E_a}{RT}$$

(6)

$$\ln \left( \frac{dm_c}{dt} \frac{1}{C_{CO_2}} \right)^2 = \ln \left( S_p^2 k_0 DS_p \right) - \frac{E_a}{RT}$$

It is important to note that Equation (5) takes into account diffusion processes under special conditions, rather than being a generalization of Equations (1) and (3). Therefore, the scope of applicability of Equation (6) to the processing of thermal analysis data is limited to pore-diffusion regimes. To the authors’ knowledge, there is no published research on the application of Equation (6) to the interpretation of thermal analysis results.

All models considered herein were used for processing kinetic curves of charcoal gasification in CO₂. The pre-exponential factor is determined by the area of the reacting surface, which is usually measured with a high degree of uncertainty. Therefore, the main kinetic characteristic for comparing different models is the activation energy. As is known from published research data, the average activation energy is of the order of 200 kJ mol⁻¹ [54,57,58]. Therefore, the gasification regime of samples under thermal analysis conditions can be identified by the proximity of the apparent activation energy to the anticipated value.

3. Results

3.1. Thermogravimetric Curves

The obtained thermogravimetric curves are shown in Figure 3. With increasing heating rate, the experiment time decreases, so at heating rates of 20 and 50 K min⁻¹, the sample does not have enough time to achieve full conversion (due to heat resistance limitations of the furnace unit). In addition, further heating can lead to ash melting, which can distort kinetic measurements. Moreover, the two stages are rather clearly
distinguishable within the curves. The first one corresponds to the volatile residual yield. The second stage corresponds to carbon gasification, which begins from 850–1000 K.

Figure 3. TG curves for charcoal gasification in the carbon dioxide atmosphere at different heating rates (1, 5, 10, 20, and 50 K min⁻¹).

3.2. Kinetic Analysis

3.2.1. External Diffusion Model

Let us consider the TG curve of charcoal gasification at a heating rate of 10 K min⁻¹. Figure 4a shows the ln(-dm/dt)–ln(T/273) plot. The correlation proves quite good, but the exponent p determined by the slope of the regression line turns to be 11.9. The anticipated value of p for the diffusion regime of conversion, as stated above, is of the order of 1–2, so the diffusion model is not suitable for describing the gasification kinetics under the conditions of the experiment conducted.

Figure 4. Results of processing of the TG curve of charcoal gasification at a heating rate of 10 K min⁻¹ using the equation of diffusion (a) and pore-diffusion kinetics (b).

3.2.2. Pore-Diffusion Model

Data processing within the framework of the pore-diffusion reaction model yields the activation energy value of 212 kJ mol⁻¹, which is quite close to the anticipated value (Figure 4b). It can be concluded that in the case of charcoal gasification given a heating
rate of 10 K min⁻¹, the selected mass of the sample, and the flow rate values, the pore-diffusion regime is observed.

3.2.3. Kinetic Model

Linearization using Equation (2) with different exponents of the surface-mass relationship \( n \) (0 to 4) also generates the seemingly satisfactory results: Correlation coefficients for all models turn out to be at least 0.9 (Figure 5). However, the estimation of activation energy at values \( n \) equal to 1 yields the value of the order of 100–130 kJ mol⁻¹, which is approximately half of the anticipated value [57]. A suitable activation energy value can be found given \( n = 4 \) (200 kJ mol⁻¹), but it is difficult to physically justify this value of \( n \).

![Figure 5](image)

Figure 5. The results of processing of the TG curve of charcoal gasification at a heating rate of 10 K min⁻¹ using Equation (2) with different exponents: (a) \( n = 0 \); (b) \( n = 1 \); (c) \( n = 2/3 \); (d) \( n = 0.5 \); (e) \( n = 2 \); (f) \( n = 4 \).

3.3. Discussion of Kinetic Analysis Results

3.3.1. Results of Experimental TG-Curves Processing

Arrhenius plots constructed for TG curves of charcoal gasification at other heating rates are presented in Figures 6–9. At a heating rate of 1 K min⁻¹, the application of kinetic model (2) at \( n = 1 \) yields the activation energy value of 197 kJ mol⁻¹ (Figure 6). Most likely, at low heating rates, the kinetic regime is observed. However, already at a heating rate of 5 K min⁻¹, a simple kinetic model (2) yields activation energy values of the order of 100–130 kJ mol⁻¹ (see Figure 7), just like in the above case with a heating rate of 10 K min⁻¹. The application of the pore-diffusion kinetics Equation (6) results in a activation energy value of 209 kJ mol⁻¹. The results of processing of TG curves for a heating rate of 20 K min⁻¹ by Equation (6) yield an activation energy value of 193 kJ mol⁻¹, which is also quite close to the anticipated value (Figure 8).
Figure 6. Results of processing of charcoal gasification TG curve at a heating rate of 1 K min⁻¹ by different models: (a) $n = 0$; (b) $n = 0.5$; (c) $n = 2/3$; (d) $n = 1$; (e) diffusion model; (f) pore-diffusion model.

Figure 7. Results of processing of the charcoal gasification TG curve at a heating rate of 5 K min⁻¹ by different models: (a) $n = 0$; (b) $n = 0.5$; (c) $n = 2/3$; (d) $n = 1$; (e) diffusion model; (f) pore-diffusion model.
At a heating rate of 50 K min\(^{-1}\) (Figure 9), there are significant deviations from the anticipated value when using any of the models. Arguably, at high heating rates, the effects of the thermal inertia of the sample may prove significant. Due to intensive endothermic reaction on the surface, temperature irregularities occur, and the use of kinetic methods requires factoring them [29].

3.3.2. Discussion

The results of the processing of TG curves obtained at different heating rates using different kinetic models are given in Table 1. As can be seen, at the minimum heating rate, the kinetic model with the parameter \( n \) equal to one yields the most adequate value of the
activation energy: At slow heating, the sample conversion is completed by about 1150 K, and the charcoal gasification, in this case, proceeds in the kinetic regime. Such low heating rates, however, are inconvenient in experimental studies as they require more time to perform the experiment. With a heating rate of 5–20 K min⁻¹, the pore-diffusion model (denoted in Table 2 as Z) produces better results. Since the experiment time is 5 and 20 less than at a heating rate of 1 K min⁻¹, the model of pore-diffusion reaction may become a standard model for processing the kinetic curves of carbon gasification in CO₂ under these conditions. At a heating rate of 50 K min⁻¹, as mentioned above, there are additional difficulties in interpreting the TG curve: This heating rate is not recommended for studies of such processes. It can be assumed that the applicability of the pore-diffusion model for determining the activation energy (according to our experimental data) is limited by heating rates in the range of 5–20 K min⁻¹.

Table 2. Activation energy values of reaction C + CO₂ (kJ mol⁻¹) based on the results of TG curves processing.

| Model (Expression) | 1   | 5   | 10  | 20  | 50  |
|---------------------|-----|-----|-----|-----|-----|
| n = 0 (2)           | 118.46 | 104.64 | 106.36 | 96.46 | 80.46 |
| n = 1/2 (2)         | 157.84 | 116.99 | 118.04 | 118.49 | 108.14 |
| n = 2/3 (2)         | 170.97 | 121.11 | 121.93 | 125.84 | 117.37 |
| n = 1 (2)           | 197.23 | 129.34 | 129.72 | 140.53 | 135.83 |
| n = 2 (2)           | 187.59 | 154.03 | 153.08 | 184.60 | 191.20 |
| Z (6)               | 236.92 | 209.28 | 212.71 | 192.92 | 160.92 |

Thus, our proposed model allows us to explain the sensitivity of the apparent activation energy to heating conditions. Applying the usual kinetic model (n = 1), we obtain a scatter of the activation energy values of 129–197 kJ/mol. Having accepted the hypothesis about the effect of pore diffusion, we obtain the values of 192–212 kJ/mol (i.e., range becomes almost three times smaller). Note that the chemical mechanism of the heterogeneous reaction does not change in this case (at least this change is not considered). Conditions of mass transfer between reacting surface and external gas flow changes only. A rigorous confirmation of this hypothesis requires more detailed measurements (first of all, with varying the characteristic size of the samples) and has not yet been performed.

Isothermal experiments on gasification of charcoals in CO₂ atmosphere are presented in the literature, for example, in works [9,45,59,60]. In order to determine the mass transfer regime in these experiments, the apparent activation energy values can be compared. In paper [9], the activation energy of the charcoal gasification reaction at temperatures above 1120 K is equal to 123 kJ/mol. The activation energy is 100–110 kJ/mol in paper [61], and is 130 kJ/mol in paper [45] (for agricultural waste char). It can be assumed that the results obtained in these works also correspond to the proposed diffusion-kinetic model. It should be noted that Equation (5) describes the reaction-diffusion interaction in quasi-stationary approximation, so reaction rate does not depend on conversion degree. Therefore, at the initial stage of conversion, the mass loss rate in isothermal conditions should be approximately constant, which is observed in a number of experimental works (for example, References [9,60]). When the pore diffusion region becomes smaller than the size of the unreacted sample, the gasification rate depends on the parameters of individual particles, and Equation (5) becomes inapplicable. In this case, to estimate the Arrhenius coefficients, expressions are required that include the dependence on the conversion degree. This circumstance can explain the tendency towards an increase in the activation energy at high temperatures in our experiments (see Figures 5 and 7–9).

At the end of the section, we are to clarify important details regarding the determination of the kinetic parameters from one kinetic curve. The guidelines [62] state the preferred use of kinetic analysis methods using multiple kinetic curves. Such an
analysis is applicable in the case when the apparent activation energy does not depend on
the heating rate. If, at different heating rates, the determining stages of the whole process
change (in our case, mass transfer and chemical reaction), kinetic analysis using several
curves obtained under different conditions will be inappropriate. This work, however,
does not assert the erroneousness of recommendations [62], but points to some limitations
of their applicability in the case of heterogeneous reactions (on the example of carbon
gasification).

4. Conclusions

This paper proposed a method for determining the kinetic coefficients of the
heterogeneous reaction (charcoal gasification by carbon dioxide) under pore-diffusion
control (the rates of heterogeneous reaction and oxidizer diffusion in the porous space of
the sample have the same order of magnitude). For TG-curves processing, a quasi-
stationary approximation is used. The pore-diffusion model makes it possible to correct
the values of the kinetic parameters, which are obtained using commonly used models
(volume reaction, shrinking core). The range of activation energy variation changes from
129–197 kJ/mol to 192–212 kJ/mol, i.e., it becomes almost three times smaller.

At the most commonly used heating rates of 5–20 K min⁻¹, there is fairly good
correspondence between the experimental curves and the model of porous-diffusion
kinetics. At a heating rate of 1 K min⁻¹, the mass transfer constraints become insignificant,
and a simple kinetic model produces satisfactory results. At a heating rate of 50 K min⁻¹
the chemical reaction takes place at higher temperatures, the average gasification rate
increases: Apparently, under such conditions, the assumed approximations (quasi-
stationarity, temperature uniformity) become inapplicable. The diffusion model is
inapplicable in all cases.

The obtained results narrow down the area of parameters to the range that enables
conducting kinetic studies under the conditions of thermal analysis and also provide tools
for determining the kinetic coefficients of charcoal gasification under the influence of
diffusion processes. The conditions for the transition of a heterogeneous reaction in the
pore-diffusion control may depend, in addition to the heating rate, on the characteristic
size of the sample and the features of gas flow. Therefore, the found conditions may
change when using other sample mass, crucible type, or gas composition. A more rigorous
determination of these dependencies is the subject of future research.

Author Contributions: Conceptualization, I.D.; methodology, I.D.; samples preparation and
experiments, A.K.; writing—original draft preparation, I.D. and A.K. All authors have read
and agreed to the published version of the manuscript.

Funding: The research was carried at the MESI SB RAS under the program for basic research of the
SB RAS (project ID FWEU-2021-0005, reg. No. AAAA-A21-121012190004-5) using the equipment of the
“High-Temperature Circuit” scientific center.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: The data used in this study are available by request to the authors.

Conflicts of Interest: The authors declare no conflict of interest.

Notation

\( m_c \) carbon mass
\( t \) time
\( k_0 \) pre-exponential factor
\( E_a \) activation energy
\( R \) universal gas constant
\( T \) temperature
S \quad \text{surface area} \\
C \quad \text{oxidizer concentration} \\
n \quad \text{exponent in the formula for the dependence of reaction surface on sample mass} \\
(N) \quad \text{reaction order with respect to fuel} \\
N_uo \quad \text{mass transfer Nusselt number} \\
D \quad \text{diffusion coefficient} \\
L \quad \text{characteristic linear crucible size} \\
p \quad \text{exponent in the formula for the dependence of the diffusion coefficient on temperature}

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