The Effect of Al Particles Size on the Thermal Behavior and Kinetics of Al-MnO₂ Thermite System

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Micron-MnO₂ powder has unique thermal decomposition process compared with other metal oxides, and the different characteristics of components in thermite could affect the thermal performance of the whole system directly. In this work, the Al powder with different three particle sizes was combined with micron-MnO₂ to prepare the Al-MnO₂ thermite system, and the effect of Al powder particle sizes on the whole thermal behavior was studied. Firstly, the thermal decomposition process of micron-MnO₂ and purity of Al powder are tested by TG-DSC. By using ultrasonic dispersion method, the fuel-rich thermite samples were prepared and characterized by SEM and TG-DSC at different heating rates. The Kissinger method was also employed to calculate the activation energy for the first exothermic peak. It was found that the thermal decomposition process of MnO₂ in the thermite system can be significantly disturbed by different Al particles size. In other words, the effect of Al particle sizes on the thermite can be magnified due to the unique decomposition properties of micron-MnO₂ instead of onset temperature of exothermic reaction changing simply. The activation energy of thermite system decreased with the reduction of Al particles sizes in micron-level, while in nanolevel the activation energy markedly increased. Finally, the possible reasons for phenomenon were discussed.

1. Introduction

Energetic materials could rapidly release enormous heats and energy during the reaction [1–3]. Thermites, as a kind of energetic materials, are broadly employed in a wide range of applications, including micropropulsion, gas generators, welding, electric ignitors, and ammunition primers because of their high adiabatic flame temperature, flame propagation velocity, and energy density [3–7]. In general, thermite systems are the mixture of an oxidizer and fuel.

Aluminum (Al), as a main fuel in thermite system, usually mixes with the other metal oxidizers to form the Al-based thermites. The most widely reported Al-based thermites are Al-Fe₂O₃, Al-MoO₃, Al-WO₃, Al-CuO, and Al-KMnO₄ [7–13]. Hu et al. [7] synthesized the pollen-like porous Al/Fe₂O₃ thermite by a template method. During differential scanning calorimetry (DSC) tests, the reactivity of the thermite is efficiently improved corresponding to its enlarged contact surface area between Al nanoparticle and the pollen-like porous Fe₂O₃. The thermal behavior of the metastable intermolecular composite composed of the Al nanoparticles and MoO₃ is studied with DSC tests as a function of the size and size distribution of the Al nanoparticles. And the results indicate that the reactivity of Al nanoparticles is significantly higher than that of the micrometer-scale samples [9]. WO₃ is of interest as an oxidant for metals in thermites. The nanoscale WO₃ is developed by wet chemistry method to make the Al/WO₃ thermite system. And the performance of the Al/WO₃ thermite system is detailed [10]. Besides, CuO is the common metal oxide added into thermite system as reductant. The thermite reaction between the CuO nanowires and the deposited nano-Al is studied. Compared with CuO and Al nanopowders mixture, the heat released of CuO nanowires
coated with nano-Al is higher, and the higher heat formation mechanism is explained by the intimate contact [11].

But based on Fischer’s research, using manganese dioxide (MnO₂) as a metal oxidizer formed the Al-MnO₂ thermite system which also has high heat of reaction theoretically [13] but lacks particular experiments and researches, especially the effect of Al particles size on the thermal behavior of Al-MnO₂ thermite system. The estimation of isothermal values of activation energy for Al-MnO₂ thermite system is reported at an early stage [14], and the molar ratio of MnO₂ and Al is about 1:5. But the molar ratio of MnO₂ and Al should be about 3:4 from the chemical equation. Recently, Kelsey and coworkers choose Mg powder as the fuel rather than Al powder to study the effects of rheological properties on the reactivity of energetic thin films of MnO₂-Mg [15]. But in terms of application security, the stability of Al powder is better than Mg powder. Namely, Al powder has a relatively lower sensitivity than Mg powder. Therefore, we are much interested in MnO₂-Al thermite system.

Although so many previous studies have shown the properties and performances of many types of thermite system, the effect of Al particles size on thermal behavior and kinetics of Al-MnO₂ thermite system has not been well documented. Besides, the thermal process of micron-MnO₂ is significantly different from other ordinary metal oxidizers, such as CuO, Fe₂O₃, MoO₃, and WO₃. Micron-MnO₂ has two processes of thermal decomposition at the range of room temperature to 900°C [16–18]. In this work, the thermite systems of different Al particle sizes mixed with micron-MnO₂ were prepared by ultrasonic dispersion method. Based on the TG-DSC tests at different heating rates, the famous Kissinger method was chosen to calculate the activation energy of the samples preliminarily at their first exothermic peaks. The objectives of this study are to figure out the changes of thermal behavior and kinetics of Al and micron-MnO₂ thermite system with different Al particles sizes.

2. Experimental

2.1. Materials and Sample Preparation. All chemicals were analytical reagent grade and were used without any further treatment or purification. The micron-MnO₂ (5 μm) and Al powders (5 and 1 μm, 100–200 nm) were purchased from Nai-ou Nano Technology Co., Ltd. (Shanghai, China). The absolute ethyl alcohol was purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China).

In this paper, the purity of Al powder should be tested by using TG-DSC under air atmosphere before preparing the thermite samples. Then, thermite mixtures of Al with different particles size and MnO₂ are prepared using ultrasonic dispersion method. In order to make full use of MnO₂ oxidizer, the fuel-rich formula was selected. The Al/MnO₂ mixture dispersed in absolute ethyl alcohol was sonicated for 60 min in a sonic bath to prepare a homogeneously dispersed solution. After sonication, the slurry was dried at 60°C for 12 h in vacuum oven.

2.2. FE-SEM and XRD Analysis. The morphology, particle size, and mixing quality of the materials and mixtures were characterized by field emission scanning electron microscopy (FE-SEM) analysis (HITACHI S-4800 II, Japan, and Zeiss EVO, Germany). The phase structures of thermite samples and the reaction products were tested by using X-ray diffraction (XRD) analysis (Bruker, D8 Advance, Germany).

2.3. Thermal and Kinetics Analysis. Investigations of thermal behavior of the pure compounds and mixtures were carried out using the TG-DSC simultaneous thermal analyzer (NETZSCH STA 449F3, Germany). As for the pure compounds, the heating rate was 10°C·min⁻¹. And as for the Al/MnO₂ mixtures, the heating rates were 8, 10, and 14°C·min⁻¹ (covering the temperature range from room temperature to 1000°C). Nitrogen atmosphere was chosen as purge and protective gas.

In this work, the famous Kissinger method was used in the calculation of thermal kinetics of the Al/MnO₂ thermite mixtures. The activation energy was calculated through Kissinger method based on the DSC peak temperature [19–21]. This method can be expressed by the following equation:

\[
\ln \left( \frac{\beta}{T_p^2} \right) = \ln \left( \frac{A}{E_a} \right) - \frac{E_a}{R T_p}
\]

where \( \beta \) is the linear heating rate (°C·min⁻¹), \( T_p \) the absolute temperature of DSC peak temperature (K), \( R \) the universal gas constant (J·mol⁻¹·K⁻¹), \( A \) the preexponential factor (s⁻¹), and \( E_a \) the activation energy (kJ·mol⁻¹). Thus, the plot of \( \ln (\beta/T_p^2) \) versus \( 1/T_p \) should be a straight line whose slope can be used to evaluate the activation energy. The degree of credibility will be higher if the absolute value of the correlation coefficient is much closer to 1 theoretically.

3. Results and Discussion

3.1. Thermal Properties of Micron-MnO₂. In order to verify the thermal decomposition of the pure 5 μm MnO₂, the TG and DSC curves of 33.564 mg MnO₂ sample into the 80 μL corundum crucibles are obtained in nitrogen atmosphere from room temperature to 900°C, as shown in Figure 1. In Figure 1, the process of thermal decomposition can be divided into two steps.

The first step of thermal decomposition takes place between 507.3°C and 598.1°C with a sharp mass loss of about 8.60% because of O₂ production. The heat release of the first decomposition step is about –197.1 Jg⁻¹. Besides, the second step of thermal decomposition takes place between 734.6°C and 842.6°C with a mass loss of about 3.07% because of O₂ production, too. The heat release of the second decomposition step is about –57.4 Jg⁻¹. According to the previous studies, the thermal decomposition of MnO₂ can be summarized as follows: [16–18],

\[
\text{MnO}_2 \xrightarrow{500–600°C} \text{Mn}_2\text{O}_3 \xrightarrow{730–850°C} \text{Mn}_3\text{O}_4
\]
3.2. Purity of Al Powder Tests. In fact, the alumina shell effect on Al powder usually cannot be negligible simply, especially for the nano-Al powder. So, in order to figure out the purity of Al rigorously, the TG-DSC tests were carried out under air conditions with 10°C·min\(^{-1}\) heating rate, as shown in Figure 2. According to the reaction equation, the Al can be oxidized by gaseous O\(_2\) leading to the Al\(_2\)O\(_3\) product, so theoretically the percentage increase in mass should be about 88.89%. However, due to the existence of alumina shell, from room temperature to 1000°C, the values of mass increase of 5 \(\mu\)m Al powder, 1 \(\mu\)m Al powder, and nano-Al powder are 57.27\%, 50.02\%, and 40.96\%, respectively. Namely, the purity of 5 \(\mu\)m Al powder is about 83.26\% and purity of 1 \(\mu\)m Al powder is about 79.42\%, while that of nano-Al powder is only about 74.62\%.

3.3. FE-SEM and XRD Analysis. Figure 3 shows FE-SEM images of the pure components. Particle shapes of 5 \(\mu\)m MnO\(_2\) are shown in Figure 3(a); the particles are irregular polygon with partial agglomeration. The average 5 \(\mu\)m Al particles are shown in Figure 3(b) while the average 1 \(\mu\)m Al particles are shown in Figure 3(c), which belong to the micron-level Al powder. As shown in Figure 3(d), it is the morphology of nano-Al powder with the average diameter of 100–200 nm. Compared with the micron-level Al powder, the nano-Al powder shows the great mass of agglomeration.

Figure 4 shows SEM images of different thermite systems. The morphologies and distribution of 5 \(\mu\)m Al + micron-MnO\(_2\) and 1 \(\mu\)m Al + micron-MnO\(_2\) thermite samples are shown in Figures 4(a) and 4(b), respectively, and the contact of 1 \(\mu\)m Al + micron-MnO\(_2\) thermite sample is more intimate than that of 5 \(\mu\)m Al + micron-MnO\(_2\) thermite sample. The nano-Al + micron-MnO\(_2\) thermite sample is shown in Figure 4(c). Since a large number of nano-Al particles attach to the surface of MnO\(_2\), the surface-area-to-volume ratio could increase a lot. Besides, the nano-Al particles still have the obvious agglomeration phenomenon.

Figure 5 shows the XRD results of thermite samples. The phases of MnO\(_2\) bulk in thermite systems are consistent with the tetragonal pyrolusite (ICDD/JCPDS 24-0735 MDI Jade 6.0). The space group is P42/mmm (136), and the lattice constants are \(a = b = 4.399\) Å, \(c = 2.874\) Å. Besides, the selected Al powders at different particle sizes have the same phase from the XRD results, which are consistent with aluminum (ICDD/JCPDS 04-0787 MDI Jade 6.0). The space group is Fm-3m (225), and the lattice constants are \(a = b = c = 4.049\) Å. The XRD results can rule out the effect of crystal shape on the thermal analysis of thermite samples.

3.4. Thermal Properties Analysis. The thermal properties of thermite samples were tested by using TG-DSC at 10°C·min\(^{-1}\) at nitrogen atmosphere, as shown in Figure 6.

Figure 6(a) represents the thermal behavior of 5 \(\mu\)m Al + micron-MnO\(_2\) thermite sample from room temperature to 1000°C. During the TG-DSC testing, there is no evident reaction or physical change from room temperature to 500°C. As the test temperature rises further, there are six DSC peaks of A, B, C, D, E, and F sequentially in Figure 6(a).

Endothermic peak A, from 512°C to 591°C and with peak temperature at 566°C, is on behalf of the first decomposition of MnO\(_2\) due to the mass loss and endothermic process, which can be judged by TG curve. The products are Mn\(_2\)O\(_3\) and gaseous O\(_2\) release leading to about 4.5% mass loss. However, according to the results of thermal decomposition of micron-MnO\(_2\) in Figure 1, if all of the micron-MnO\(_2\) in the thermite system has taken part in the decomposition, the loss of mass in this temperature interval should be apparently more than 4.7%. Namely, some of MnO\(_2\) does not decompose at all but reacts with 5 \(\mu\)m Al powder with releasing a little heat at the same time, which should be exothermic peak B. The peak temperature of peak B is 594°C with only 57 Jg\(^{-1}\) heat release. As for exothermic peak C, there is a clear gap between peaks B and C, indicating that peak C is on behalf of different reaction process. It should be the reaction between 5 \(\mu\)m Al powder and products after decomposition of MnO\(_2\), part of Mn\(_2\)O\(_3\) and part of gaseous O\(_2\) due to the mass increase a little, and the peak temperature is 626°C. The above exothermic reactions appear before the Al powder melt, which belongs to solid-phase reaction. Endothermic peak D means the melt of Al at about 660°C. Then, endothermic peak E appears with the mass loss further in TG curve, which means the second thermal decomposition of MnO\(_3\) from remaining Mn\(_2\)O\(_3\) to MnO\(_4\) with gaseous O\(_2\) release. Next, at the temperature range 850°C–960°C, there is the main exothermic peak F, which is thermite reaction between molten Al and Mn\(_2\)O\(_4\) as well as part of generated O\(_2\). The peak temperature of peak F is at 943°C. In summary, the 5 \(\mu\)m Al + micron-MnO\(_2\) thermite system shows the low reactivity, which might be because the distance between the fuel and oxidizer is not close enough from SEM image.
rises further, there are five DSC peaks of A, B, C, D, and E sequentially in Figure 6(b). Compared with the peaks of 5 μm Al + micron-MnO₂ thermite sample in Figure 6(a), the first four peaks, A, B, C, and D, are similar, indicating the same processes. Endothermic peak A, from 475°C to 593°C, means the first thermal decomposition of MnO₂. Exothermic peak B represents the thermite reaction between MnO₂ and 1 μm Al powder while exothermic peak C is the redox reaction between 1 μm Al powder and part of Mn₂O₃ as well as part of gaseous O₂. But the difference is that more MnO₂ is involved in the Al-MnO₂ thermite reaction directly judged from the more heat release from peak B, about 193 J g⁻¹ and less mass loss in TG curve, only about 4.1%. Undoubtedly, endothermic peak D is the melting process of Al powder. However, as for exothermic peak E in Figure 6(b), it is the main difference from Figure 6(a). Both DSC curve and TG curve show that the further thermal decomposition does not happen. There is not any endothermic signal or mass loss signal. There is only an obvious exothermic peak E at the temperature range 788°C to 882°C, and the peak temperature of peak E is at 833°C in advance. Namely, all of the remaining MnO₂ reacts with the molten Al directly with 418 J g⁻¹ heat release. So, if the Al powder particle sizes decrease to 1 μm, the reactivity of the whole thermite system increases greatly. More MnO₂ reacts with Al powder directly, and all of remaining Mn₂O₃ reacts with molten Al rather than the further decomposition.

In Figure 6(c), the nano-Al powder is introduced to the Al-MnO₂ thermite system. At the beginning of the TG-DSC, there is a slight mass loss in the TG curve, about 0.89%, from room temperature to 300°C without any evident DSC signal, which is due to the desorption of H₂O and ethyl alcohol on the surface of nano-Al powder [22, 23]. In contrast, there are merely four peaks in DSC curve, peaks A, B, C, and D. As for
peak A, it becomes an exothermic peak rather than an endothermic peak in Figures 6(a) or 6(b). The heat release reaches about 735 J g⁻¹, and the temperature range is from 503°C to 591°C. At the same time, as for TG curve, no significant mass loss or mass gain can be found at the range of temperature from 300°C to 600°C, namely, all of MnO₂ takes part in the thermite reaction at solid-phase state rather than the first thermal decomposition. The thermite reaction, meaning oxygen transfer from the MnO₂ to the Al, should not exhibit any change in the mass signal due to conservation of mass. No other mass loss can be found as temperature rises further, which means that all of the MnO₂ is consumed in the first main thermite reaction. Then, after the thermite reaction, exothermic peak A, there are still three

![Figure 3: SEM images for the components: (a) micron-MnO₂, (b) 5μm Al powder, (c) 1μm Al powder, and (d) nano-Al powder.](image)

![Figure 4: SEM images of thermite samples: (a) 5μm Al + micron-MnO₂ thermite sample, (b) 1μm Al + micron-MnO₂ thermite sample, and (c) nano-Al + micron-MnO₂ thermite sample.](image)
Figure 5: XRD pattern of the thermite samples.

Figure 6: TG-DSC curves of thermite samples with different Al particle sizes: (a) 5 μm Al + micron-MnO₂ thermite sample, (b) 1 μm Al + micron-MnO₂ thermite sample, and (c) nano-Al + micron-MnO₂ thermite sample.
Figure 7: XRD pattern of the reaction products.

Figure 8: Continued.
peaks, exothermic peaks B and D and endothermic peak C. Clearly, endothermic peak C is due to the melting of Al powder. As for exothermic peaks B and D, they should be the reaction between nano-Al and N$_2$ with the product of aluminum nitride (AlN) since the obvious mass gain in the TG curve. Since the TG-DSC is running in the nitrogen atmosphere and the mass gain is due to the reaction of the Al to nitrogen in the environment, AlN is produced, releasing part of the heat [24].

At the end of the TG-DSC tests, the residues in the crucibles are collected, and the reaction products are tested by using XRD analysis, as shown in Figure 7. The phases of reaction production in 5μm Al + micron-MnO$_2$ thermite sample are mainly Mn$_3$O$_4$ and part of MnO. When the particle size of Al powder decreases to 1μm, the main reaction products still are Mn$_3$O$_4$ and MnO, but the mass fraction of MnO in residues increases. Based on the results of thermal analysis in Figure 6, a certain part of Mn$_3$O$_4$ in the residues of 5μm Al + micron-MnO$_2$ thermite sample should decompose from the Mn$_3$O$_7$ while most of Mn$_3$O$_4$ in the residues of 1μm Al + micron-MnO$_2$ thermite sample could be the reaction products from the thermite reaction between Mn$_3$O$_7$ and molten Al, which also reflects the different reactivity between the above two thermite samples. As for nano-Al + micron-MnO$_2$ thermite sample, the residues are mainly MnO with only little Mn$_3$O$_4$. Namely, nano-Al powder has much more reactivity, so it could make full use of the O element in the oxidizer MnO$_2$.

3.5. Activation Energy. In order to make a further comparison of kinetics, the thermite samples were also tested by DSC at different heating rates. The thermite has a phenomenon of self-propagating combustion once it ignites successfully, namely, the first exothermic peak temperature is important in practice, as shown in Figure 8, and the peak temperature points are listed in Table 1.

As seen in Table 1, in terms of peak temperature for the different sample components, the peak temperature decreases with the decrease of particle size of Al powder. When Al particle size enters nanolevel, peak temperature has a significant reduction, near 560°C. However, the average heat release is the opposite of peak temperature. Namely, the average heat release increases with the decrease of particle size of Al powder.

The Kissinger method is one of the dominant methods in the maximum rate methods (peak methods) [25, 26]. Based upon Kissinger method mentioned in equation (1), the plots of ln ($\beta / T_p^2$) vs. $1/T_p$ at the peak temperature with the different Al particles size are constructed in Figure 9.
absolute values of all correlation coefficient $R$ were above 0.94. And the figures of activation energy $E_a$ were listed into Table 2, which were deduced from the slope of the data correlation line. According to Table 2, when the Al particle size decreased from 5 μm to 1 μm, the activation energy of the thermite decreased. But if the Al particle size decreased further into the nanolevel, the activation energy of the thermite increased markedly. As known to all, the Al powder has an oxide layer coating on the surface. Usually, the thickness of oxide layer can greatly reduce the purity of nano-Al powder, which is corresponding with the results of purity tests in Figure 2. Namely, as for micron-level Al powder, the thickness of oxide layer has a very light proportion for Al powder [26, 27]. The reduction of reactive Al purity in nano-Al particles could be one of the reasons for activation energy increase. Besides, the agglomeration of

| Thermite formula          | 5 μm Al micron-MnO$_2$ | 1 μm Al micron-MnO$_2$ | Nano-Al micron-MnO$_2$ |
|---------------------------|------------------------|------------------------|------------------------|
| $E_a$                     | 283.76 kJmol$^{-1}$    | 153.90 kJmol$^{-1}$    | 415.48 kJmol$^{-1}$    |

Figure 9: The data correlation line of ln (β/1/T$^2$) vs. 1/T$_p$: (a) 5 μm Al + micron-MnO$_2$ thermite, (b) 1 μm Al + micron-MnO$_2$ thermite, and (c) nano-Al + micron-MnO$_2$ thermite.
nano-Al particles cannot be ignored. Agglomeration could be very important for the characteristic of nano-level Al powder, such as ignition kinetics and heat diffusion. The large size agglomeration might ignite after very long delays or even never ignite during the whole combustion process [28]. Thus, the agglomeration will not only decrease the heat of reaction, but also greatly increase activation energy of thermite system [29–31]. So, the above two reasons could explain the phenomenon of the significant increase of activation energy of nano-Al+micron-MnO2 thermite sample.

4. Conclusion

In this paper, Al-MnO2 thermite samples with three different kinds of Al particle sizes were prepared via ultrasonic dispersion method. Firstly, the thermal decomposition process of micron-MnO2 and the purity of Al powders were tested by using TG-DSC. The values of purity of 5 μm Al powder, 1 μm Al powder, and nano-Al powder were 83.26%, 79.42%, and 74.62%. According to XRD analysis, the components in the thermite system were same, and the only difference was the particle sizes of fuel, Al powders, from the SEM images. On the basis of that, the fuel-rich thermite systems were tested for their thermal properties and performance. The results showed that 1 μm Al powder + micron-MnO2 thermite system has better reactivity than that of 5 μm Al powder + micron-MnO2 thermite system. The latter has six DSC peaks while the former has only five DSC peaks without the second endothermic peak of thermal decomposition. However, as for nano-Al powder + micron-MnO2 system, there are only four DSC peaks. Apart from the melting of Al powder, there is not any other endothermic peak. Namely, almost all of micron-MnO2 reacted with solid-state nano-Al powder directly. After collecting and testing the residues, the reaction products of 5 μm Al powder + micron-MnO2 thermite system were mainly Mn3O4 and a little MnO while that of 1 μm Al powder + micron-MnO2 thermite system showed a bit more fraction of MnO, but as for nano-Al powder + micron-MnO2 thermite system, the residues were mainly MnO with little Mn3O4, which reflected that nano-Al made full use of O element in oxidizer MnO2. However, the kinetics results showed that nano-Al powder + micron-MnO2 thermite system had the highest activation energy, which can be explained by low reactive Al purity and the agglomeration of Al nanoparticles. The results clearly showed the great different thermal processes and changes of Al-MnO2 thermite samples with different particle sizes of Al powder.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest

The authors declare that there are no conflicts of interest regarding the publication of this paper.

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