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Enhancing Aluminum Reactivity by Exploiting Surface Chemistry and Mechanical Properties

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Many Al fueled energetic composites use solid oxidizers that induce no alumina surface exothermicity, such as molybdenum trioxide (MoO3) or copper oxide (CuO). To further understand the role of PFPE in energetic systems, varying concentrations were blended with Al/CuO and Al/MoO3 thermites. Flame speeds, differential scanning calorimetry (DSC) and quadruple mass spectrometry (QMS) were performed for varying percent PFPE blended with Al/MoO3 or Al/CuO in order to examine reaction kinetics and combustion performance. X-ray photoelectron spectroscopy (XPS) was performed to identify product species. Results show that the performance of the thermite-PFPE blends is highly dependent on the bond dissociation energy of the metal oxide. Fluorine-aluminum based surface exothermic chemistry with MoO3 produce an increase in reactivity while the blends with CuO show a decline when increasing the PFPE loadings. These results provide new evidence that optimizing aluminum combustion can be achieved through activating exothermic Al surface chemistry that produces aluminum fluoride.

Another avenue for increasing Al reactivity is to alter its mechanical properties. In bulk material processing, annealing and quenching metals such as Al can relieve residual stress and improve mechanical properties. On a single particle level, affecting mechanical properties may also affect Al particle reactivity. Aluminum particles underwent thermal treatment in order to examine the effect of annealing and quenching on the strain of Al particles and the corresponding reactivity of Al and CuO composites. Synchrotron X-ray diffraction (XRD) analysis of the particles reveals the thermal treatment increased the dilatational strain of the aluminum-core, alumina-shell particles. Flame propagation experiments also show thermal treatments effect reactivity when combined with CuO. An effective annealing/quenching treatment for increasing aluminum reactivity was identified. These results show that altering the mechanical properties of Al particles affects their reactivity.

Enhancing Aluminum Reactivity by Exploiting Surface Chemistry and Mechanical

Properties

by

Jena McCollum, B.S.M.E.

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Michelle Pantoya Chair of Committee

Jharna Chaudhuri

Gordon Christopher

Stephen Ekwaro-Osire

Luciano Castillo

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ABSTRACT

Metal-fluoropolymer based systems have drawn quite a bit of attention in the combustion community in recent years due to the exothermic surface chemistry between fluorine (F) and the alumina (Al₂O₃) shell surrounding aluminum (Al) fuel particles promotes aluminum reactivity. Incorporating a liquid fluorinated oligomer, specifically perfluoropolyether (PFPE), exhibits this surface chemistry while increasing the proximity of fuel and oxidizer.

Flame speeds, differential scanning calorimetry (DSC), thermogravimetric analysis (TGA) and quadruple mass spectrometry (QMS) were performed for a variety of Al particle sizes blended with PFPE. The results show that the combustion performance of these blends is highly dependent on the Al₂O₃ concentration and exposed surface area. As Al particle diameter increases from 80 to 120 nm, the Al-PFPE blends exhibit an increase in flame speeds by 48%, but from 120 to 5500 nm, flame speeds decrease by 93%. There is a balance to optimizing Al particle reactivity with PFPE coating between activating Al particles with exothermic surface chemistry versus the unreacted alumina that contributes a thermal heat sink during energy generation.

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Another avenue for increasing Al reactivity is to alter its mechanical properties. In bulk material processing, annealing and quenching metals such as Al can relieve residual stress and improve mechanical properties. On a single particle level, affecting mechanical properties may also affect Al particle reactivity. Aluminum particles underwent thermal treatment in order to examine the effect of annealing and quenching on the strain of Al particles and the

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CHAPTER I: Fundamentals of Combustion

From early times, combustion has been observed and studied by curious minds. One of the earliest concepts was the phlogiston theory which was conceptualized by Johann Joachim Becher in the 17th century. This basically stated that all substances were comprised of three kinds of earth; the vitrifiable, the mercurial, and the combustible.

Combustion can be studied in gas (i.e., natural gas), liquid (i.e., gasoline, kerosene) and solid (i.e., coal, thermites, etc.) phases. Although the reaction mechanisms vary among the different phases, basic combustion characteristics can be calculated for any combustible medium.

Activation Energy and Heat of Combustion

Combustion reactions occur by exothermic reactions. This occurs when the energy of the products is lower than the energy of the reactants. This means that heat was released from the system. Figure 1 shows how the energy changes in these systems during the reaction.



Figure 1. Energy change of an exothermic reaction

In order to have a reaction, the energy level of the reactants must be raised from their equilibrium state to the minimum energy for the reaction to occur. The minimum energy needed to start a reaction is called the activation energy (E_a). The activation energy is related to the reaction rate (k(t)) and shown in Eq. (1) where *A* is the pre-exponential factor, *R* is the universal gas constant and *T* is the temperature.

$$\mathbf{k}(\mathbf{t}) = \mathbf{Aexp}(-\frac{\mathbf{E}_a}{\mathbf{RT}})$$
(1)

The heat of combustion (ΔH_c) is given by Hess' Law shown in Eq. (2).

$$\Delta H_{c} = \sum_{react} H_{f}^{o} - \sum_{prod} H_{f}^{o} \qquad (2)$$

This means that the ΔH_c is the difference in the heat of formation of the reactants and the heat of formation of the products. This translates to the amount of heat liberated from the reaction. The necessary information about reactants (fuel and oxidizer) and the products can be obtained in the reaction equation.

Reaction Equations

Combustion takes place by an oxidative reaction. The reactant that is reduced in the reaction is called the fuel. In gas and liquid combustion, these fuels are typically hydrocarbons. Their reduction by air or pure oxygen results in water (H₂O), carbon dioxide (CO₂), and other intermediates (CO, OH, C, O₂, N₂, etc.). In simplified combustion, these intermediates may not be present in the reaction formula. For example, the reaction equation for general hydrocarbon (C_xH_y) combustion with oxygen (O₂) is shown in Eq. (3).

$$aC_xH_y + bO_2 \rightarrow cCO_2 + dH_2O$$
 (3)

This equation assumes no production of intermediates (i.e. CO, OH, etc). The coefficients *a*, *b*, *c*, and *d* can be deduced by performing a molar balance of all elements. Because this reaction is considered for only 1 mole of C_xH_y , and C_xH_y has *ax* carbon (C) atoms, *c* must equal *ax*. Next, the hydrogen (H) atoms will be balanced. The hydrocarbon has *ay* H atoms, which means that *d* must equal

ay/2. Now that *c* and *d* have been obtained, *b* can be found. With *c*=*ax* and d=ay/2, the products contain a total of 2a(x+y/4) O atoms. This means that *b* must equal a(x+y/4). The final equation is shown in Eq. (4).

$$aC_{x}H_{y} + a(x+y/4)O_{2} \rightarrow axCO_{2} + a(y/2)H_{2}O$$
(4)

Stoichiometry

An important factor in combustion is the relationship between fuel and oxidizer in the reaction. If the reaction is fuel rich, there will be remaining fuel in the products. If the reaction is fuel lean, the products will contain unused oxidizer (i.e., O₂, N₂, etc). This characteristic is known as an equivalence ratio (ER). For

ER=1, the mixture is stoichiometric,

ER>1, the mixture is fuel rich

ER<1, the mixture is fuel lean

The equivalence ratio is calculated by comparing the actual number of moles of fuel-to-oxidizer ratio to the stoichiometric fuel-to-oxidizer ratio shown in Eq. (5) where *F* is the number of moles of fuel and *A* is the number of moles of oxidizer.

$$\mathbf{ER} = \frac{\frac{F}{A_{\text{actual}}}}{\frac{F}{A_{\text{stoich}}}} \tag{5}$$

For example, the general hydrocarbon case shown above would yield an equivalence ratio shown in Eq. (6).

$$ER = \frac{\frac{a}{(x+(\frac{y}{2})_{actual}}}{\frac{e}{(x+(\frac{y}{2})_{stoich}}} = 1 \text{ for } a = e \quad (6)$$

Equation (6) shows that for a=e, the mixture is stoichiometric. The mixture will be fuel lean is a < e and fuel rich if a > e.

Although hydrocarbon combustion is used for the example here, the same principles apply to the solid combustion that is presented in this work. Maintaining stoichiometry is critical when varying other parameters so to not change the fundamental combustion.

Aluminum Combustion

The fuel of interest in this work is aluminum (AI). Aluminum combustion is widely studied due to its high heats of combustion and energy density. The energy density of metal fuels is much higher than that of monomolecular explosives (meaning that they have the fuel and oxidizer in the same molecule (i.e., TNT)). The oxidative reaction that takes place converts the metal (Al) to the metal oxide (Al_2O_3). Aluminum particles are spherical and they contain a native oxide layer that is typically 3-5 nm in thickness.



Figure 2. Transmission electron spectroscopy (TEM) image of a nano-scale Al particle to show alumina shell thickness. The average thickness is 4.01 nm. This thickness was used in the purity calculations for micron-scale Al particles (< 1% by mass oxide).

Aluminum is pyrophoric, so this passivation layer stabilizes the particles and

allows them to be handled in oxygen-rich and moisture containing environments.

When Al is blended with metal oxides and heat is applied, Al receives

oxygen from the metal oxide and results in aluminum oxide (also known as

alumina or Al_2O_3). This reaction is shown in the simplified chemical equation shown in Eq. (7).

 $2AI + 3CuO \rightarrow AI_2O_3 + 3Cu \quad (7)$

Aluminum is also very reactive with fluorine. Fluoropolymers (i.e., polytetrafluoroethelyne (PTFE)) can be used as an oxidizer and the product expected is aluminum fluoride (AIF₃). A simplified version of this reaction is shown in Eq. (8).

$$4AI + 3C_2F_4 \rightarrow 4AIF_3 + 6C \quad (8)$$

Perfluoropolyether (PFPE)

Perfluoropolyether (PFPE) is a lubricant that is used for high performance automotive and aerospace applications due to its ability to withstand high temperatures (< 316 °C). Unless in the presence of a degrading metal (i.e., Al), PFPE is chemically inert. The chemical structure of PFPE is shown in Fig. 3.



Figure 3. The chemical structure of PFPE.

Unlike PTFE which is a fluoropolymer, PFPE is a fluorooligomer which is a fluoropolymer intermediate consisting of only a few structural units. This allows PFPE to have different molecular weights based on chain length.

Pre-ignition Reaction

For Al-based thermites, when fluorine replaces oxygen as the reducing agent, aluminum fluoride (AIF₃) is formed instead of Al₂O₃. Both AlF₃ and Al₂O₃ are comparably thermodynamically stable with heats of formation of 1510 kJ/mol and 1676 kJ/mol, respectively. However, AlF₃ formation is preferred to Al₂O₃, because Al-F (664±6 kJ/mol)(Cottrell 1958) bond formation is stronger than the Al-O bond (512±4 kJ/mol)(Cottrell 1958). In fact, F has been shown to react with the alumina passivation shell surrounding an Al particle to form AlF₃(Osborne and Pantoya 2007). The exothermic surface reaction precedes the main Al oxidation reaction and has been coined a pre-ignition reaction (PIR)(Osborne and Pantoya 2007). Figure 4 shows a DSC curve for an Al-PTFE reaction.



Figure 4. Heat flow and mass loss curves for the AI-PTFE reaction. Note the exotherm that occurs at the decomposition temperature of PTFE.

The PIR is a small but prominent exothermic peak that coincides with the decomposition temperature of PTFE and occurs before the oxidation of the AI core.

Metal Oxides

When Al particles react with metal oxides (i.e., molybdenum trioxide (MoO₃) or copper oxide (CuO)), the reaction is highly exothermic with flame temperatures exceeding 2000 K(Fischer and Grubelich 1998). This occurs because aluminum oxide has a higher heat of formation than other metal oxides (J. J. Moore and Feng 1995). Like Al, CuO has a spherical morphology but MoO₃ particles are flake structures. The differences in morphology are shown in Fig. 5.



Figure 5. SEM images of Al with (a) MoO_3 and (b) CuO.

These highly reactive blends of metallic fuel and metal oxide are called thermites. Their applications range from material synthesis(Shigeta and Watanabe 2010; Deevi, Sikka, and Swindeman 1997; Yeh and Huang 2011) and alloying(C. Crane, Pantoya, and Dunn 2010; L. L. Wang, Munir, and Maximov 1993) to joining and welding(Iordachescu et al. 2010; Chen et al. 2006).

CHAPTER II: Activating Aluminum Particle Reactivity with Fluorine Oligomer Surface Coating

Abstract

Exothermic surface chemistry between fluorine and the alumina (Al2O3) shell surrounding aluminum (AI) fuel particles promotes aluminum reactivity. This study investigates the reactivity of AI when coated with a liquid fluorinated oligomer, specifically perfluoropolyether (PFPE). Flame speeds, differential scanning calorimetry (DSC), thermogravimetric analysis (TGA) and guadruple mass spectrometry (QMS) were performed for AI- PFPE blends with varying AI particle sizes (i.e., 80, 100, 120 and 5500 nm average diameter). The results show that the combustion performance of these blends is highly dependent on the Al2O3 concentration and exposed surface area. As Al particle diameter increases from 80 to 120 nm, the AI-PFPE blends exhibit an increase in flame speeds by 48%, but from 120 to 5500 nm, flame speeds decrease by 93%. There is a balance to optimizing Al particle reactivity with PFPE coating between activating AI particles with exothermic surface chemistry versus the unreacted alumina that contributes a thermal heat sink during energy generation. These results introduce a new and simplified synthesis approach for activating Al

particle surface chemistry to promote overall reactivity, and can be useful for many energy generation applications that involve aluminum combustion.

Introduction

Energetic blends of aluminum (AI) and fluorinated oxidizers such as polytetrafluoroethylene (PTFE) have been widely studied due to the unique surface exothermic chemistry between fluorine (F) and the alumina shell surrounding the Al particles which promotes Al reactivity. Fluorine is more electronegative than oxygen ($\chi_{p,oxygen}$ =3.4 and $\chi_{p,fluorine}$ =4.0)(Atkins and de Paula 2010), and the AI-F bond is one of the strongest in nature (i.e. 664±6 kJ/mol(Cottrell 1958)). When Al reacts with F, AlF₃ is formed. The exothermic reaction has been observed by differential scanning calorimetry and associated with the identification of a small but prominent exothermic peak prior to the primary reaction(Osborne and Pantoya 2007; M. L. Pantoya and Dean 2009; Travis. R. Sippel, Son, and Groven 2013; Travis R. Sippel, Son, and Groven 2013; Miller et al. 2013; Farley et al. 2013). This peak has been coined a pre-ignition reaction (PIR) whose onset temperature is dependent on particle size (M. L. Pantoya and Dean 2009), fluorine source (K. S. Kappagantula et al. 2012; Farley et al. 2013) and equivalence ratio (Miller et al. 2013).

Typically the Al oxide shell is considered dead weight in a thermite reaction because it acts as a heat sink. By adding a fluorinated component, (such as PTFE) the shell undergoes exothermic surface chemistry which improves Al reactivity and contributes to the overall calorific output of the reaction (Osborne and Pantoya 2007; K. Kappagantula, Pantoya, and Hunt 2012; M. L. Pantoya and Dean 2009; Losada and Chaudhuri 2010; Conner and Dlott 2012). Solid PTFE introduces a limit to the fuel-oxidizer contact surface area which limits the combustion performance in these diffusion controlled reactions(K. S. Kappagantula et al. 2012). In an attempt to resolve the limitation caused by fueloxidizer contact surface area, Kappagantula et al(K. S. Kappagantula et al. 2012) investigated the combustion of Al particles with surface functionalized selfassembled monolayers (SAM). The SAM consisted of perfluorotetradecanoic (PFTD) acid bonded to the aluminum particle's native oxide shell (Al_2O_3). Functionalizing Al_2O_3 with SAM is well documented with carboxylic(Oberg et al. 2001; M. Lee et al. 2007; Karaman, Antelmi, and Pashley 2001) and fluorinated acids(K. S. Kappagantula et al. 2012). This method involves chemically attaching acid chains to the surface hydroxyl layer on the alumina surface. The hydroxyl layer on Al_2O_3 is well studied and occurs naturally under standard atmospheric conditions(Peri 1965; Knoezinger 1978; Chupas and Grey 2004; Sarbak 1997).

Aluminum functionalized with PFTD successfully increased the reactivity of Al particles combined with MoO₃. In fact, the Al-PFTD composite particle combined with MoO₃ demonstrated 86% greater flame speeds than Al+MoO₃ for the same Al and MoO₃ particle size. Kinetic analysis using differential scanning calorimetry (DSC) revealed the Al-F pre-ignition reaction (PIR) for the functionalized Al particles not present in the Al without functionalization(Travis R. Sippel, Son, and Groven 2013; K. S. Kappagantula et al. 2012; Miller et al. 2013; Farley et al. 2013). The PIR induced an earlier onset of the main exothermic reaction, thereby promoting Al particle reactivity.

A simplistic approach for coating the Al surface is to introduce a liquid phase fluorooligomer that wets the Al particle surface. Miller et al. (Miller et al. 2013) studied this idea using a fluorinated oligomer to wet the surface of Al particles. The motivation was to suspend nano-scale Al particles in a structural epoxy-based matrix. To extend the shelf life of the Al particles, they applied PFPE as a protective coating because of its thermal stability at low temperatures (up to 316 °C in an oxygen-rich environment) and potential to produce exothermic activity beyond 316 °C (Miller et al. 2013).They studied the exothermic behavior for Al-perfluoropolyether (PFPE) blends of varying weight percents using differential scanning calorimetry (DSC) and found two exothermic peaks, one of which occurred at the PFPE decomposition temperature of 316 °C. This first peak was identified as a pre-ignition reaction (i.e., PIR). They also found that the weight percent that produced the highest heat of combustion was 30% Al 70% PFPE corresponding to an equivalence ratio (ER) of 1.2 (i.e., slightly fuel rich).

Dean et al. analyzed DSC data for different Al₂O₃ particle sizes with PTFE in order to understand the role of exposed surface area on the exothermic Al-F reaction(M. L. Pantoya and Dean 2009). They found that the onset and magnitude of the PIR varied with particle size and exposed surface area for Al₂O₃ average particle sizes of 15, 30, 40 and 50 nm. They observed a decrease in PIR magnitude as Al₂O₃ particle size decreased from 50 to 30 nm and an increase in magnitude at 15 nm. Their results suggest an optimum Al particle diameter for promoting the Al-F surface exothermic chemistry is about 50 nm.

All of these studies(Osborne and Pantoya 2007; M. L. Pantoya and Dean 2009; Travis R. Sippel, Son, and Groven 2013; K. S. Kappagantula et al. 2012; K. S. Kappagantula, Pantoya, and Horn 2013; Farley et al. 2013) highlight Al-F reactions are: (1) exothermic; (2) a function of exposed Al₂O₃ surface area; (3) a function of the fluorine source (i.e., PFTD, PTFE); and (4) can activate Al reactivity with earlier onset of the main exothermic reaction. The objective of this study is to synthesize Al-PFPE formulations on a larger scale and analyze the macroscopic combustion

characteristics of the AI-PFPE reaction. This study extends the work by Miller et al. (Miller et al. 2013) who examined small (i.e. 5 mg) AI-PFPE samples for thermal equilibrium studies. All blends were prepared for a constant ER (φ =1.2). Flame speeds, DSC, thermogravimetric analysis (TGA), and quadruple mass spectrometry (QMS) was performed and analyzed for all blends in order to characterize combustion dynamics and reactivity.

Materials

Four different AI particle sizes were used and summarized in Table 1. Images obtained for 80, 100 and 120 nm AI particles by transmission electron microscopy (TEM) are shown in Fig. 6.



Figure 6. TEM representative for AI particles with average diameters of (a) 80 nm, (b) 100 nm, (c) 120 nm.

Each particle was passivated by an approximately 3.7 nm thick alumina shell as characterized by TEM analysis (Fig. 6). Due to thickness limitations with

TEM, the micron-scale particles could not be measured in this way. Instead, their size was examined using an AccuSizer 780 Optical particle size analyzer (Santa Barbara, CA). The micron-scale Al was suspended in filtered water (2mg Al/80 ml water) and sonicated for 60 minutes in cycles of 10 seconds on, 10 seconds off in order to avoid heating. The sample was then pulled using a syringe pump into the size analyzer for measurement. Particle length (diameter) is calculated and plotted as a count-based measurement (i.e., number of particles with a given diameter, in μ m units). This plot is shown in Fig. 7 and the size data is given in Table 1.



Figure 7. Particle size distribution for the micron-scale AI particles represented in number of particles with x diameter (i.e., counts).
Because the oxide layer thickness is independent of Al particle size, the oxide thickness used for purity calculations was estimated at 3.7 nm (i.e., the average of the other 3 samples).

The 80 and 120 nm average diameter AI particles were procured from Nanotechnologies (Houston, TX), the 100 nm average diameter AI was from U.S. Nanomaterials (Houston, TX) and the 5500 nm average diameter AI from Alfa Aesar (Ward Hill, MA). The PFPE (Fomblin[®] Y LVAC 25/6, average molecular weight of 3300 g mol⁻¹) was procured from Sigma Aldrich (St. Louis, MO).

Experimental Section

Equivalence ratio (ER) of 1.2 (i.e., slightly fuel rich) was maintained for all samples. An equivalence ratio is the balance for fuel (i.e. Al) and oxidizer (i.e. PFPE) in an energetic blend as shown in Eq. (9) where *M* is mass and *fuel* is Al, and *oxidizer* is PFPE.

$$ER = \frac{\left[\frac{M_{fuel}}{M_{oxidizer}}\right]_{actual}}{\left[\frac{M_{fuel}}{M_{oxidizer}}\right]_{stoichiometric}}$$
(9)

When the reaction completes with no remaining fuel (AI) or oxidizer (PFPE) in the products, it is stoichiometric (i.e., ER=1). The reaction of AI and PFPE is

shown in Eq. (10) where *E* represents other products of the reaction and *a*, *b* and *c* are the constants used to balance the reaction.

 $aCF_3O-[-CF(CF_3)CF_2O-]_x(-CF_2O-)_yCF_3 + bAI \rightarrow cAIF_3 + E$ (10)

The reaction was balanced in this manner for all particle sizes to account for the Al and Al_2O_3 respective concentrations. The alumina concentrations range from 23 to 36 mass percent for 120 to 80 nm respectively and less than 1 mass percent for the micron-scale Al particles.

Miller et al. prepared the AI-PFPE blends by hand mixing with a spatula for five minutes(Miller et al. 2013). Due to the small sample size required for DSC analysis (i.e., 5 mg), this mixing method is sufficient. However, the level of PFPE dispersion needed for analyzing energy propagation of larger quantities (i.e., 300 mg) required a modified mixing technique. A fluorinated solvent for mixing was included here which improved dispersion needed for energy propagation analysis. The fluorine in the solvent disperses the fluorine in the PFPE and suspends the AI particles to promote better coating. The solvent used in this study was Perfluorosolv-2 (PFS-2) from SPI supplies (West Chester, PA). In order to flow into the sample holders, 100 ml of solvent was used per 300 mg of Al-PFPE. The solvent, AI and PFPE were mixed in a Thinky (Tokyo, Japan) planetary mixer for 3 minutes at 1500 RPM.

Flame Speed Experiments

Due to the dramatic change in Al particle surface area between sizes, the powders reclaim with a different consistency. In order to mitigate this, an acrylic apparatus was machined with two notches (see Fig. 8).



Figure 8. Schematic of the acrylic sample holder used for drying and igniting Al-PFPE blends and time-stamped representative still frames to illustrate Al-PFPE burning behavior.

The large notch was designed to capture the excess solvent. While drying, the AI particles and PFPE settle into the smaller sub-notch. The sample mass was monitored until the solvent mass evaporated. Once dry, the bulk density of the samples ranged between 1.78-2.00 g/cm³. Percent theoretical maximum density (TMD) was calculated for each sample and is shown in Table 1.

name	average	oxide	particle	TMD	Bulk	%TMD	Specific
	particle	layer	mass		Density		Surface
	size	thickness	alumina				Area
(nm)	(nm)	(nm)	(%)	(g/cm ³)	(g/cm ³)	(%)	(1/cm)
80	76	3.9	36%	2.3	2.00	87	7.9x10 ⁵
100	91	3.6	29%	2.2	1.96	90	6.7x10 ⁵
120	121	3.6	23%	2.2	1.93	95	5.0x10 ⁵
5500	5800	3.7	<1%	2.1	1.78	96	1.2x10 ⁴

Table 1. Material Specifications for Al Fuel Particles.

For flame propagation analysis, one end of the open notch was fitted with a nickel-chromium wire for ignition. Each sample contained 300 ± 10 mg of Al-PFPE blend. The trays were loaded into a blast chamber which was fitted with an acrylic viewing window. The experimental setup is shown in Fig. 9.



Figure 9. Ignition setup for tracking energy propagation.

The AI-PFPE blends were ignited by hot wire and the flame propagated along the length of the tray. The reaction was captured using a Phantom v7 (Vision Research, Inc., Wayne, NJ, USA) fitted with a Nikon AF Nikkor 52 mm 1:2.8 lens with an F-stop of 32. The camera was placed perpendicular to the reaction path and collected data with a resolution of 800 x 512 pixels at 1000 frames per second. The reaction is captured as a series of images which are then analyzed using NI Vision Builder to obtain the change in position of the flame front as a function of time. Each reported flame speed is the average of five experiments and the uncertainty in the measurement is represented by the bars in Fig. 10. The uncertainty is <10% such that this method was successful in producing macroscopically dispersed AI-PFPE mixtures in terms of repeatable energy propagation behavior and measurements.

Equilibrium Analysis

Simultaneous Thermal Analysis (STA) was performed using a Netzsch Jupiter STA 449 differential scanning calorimeter and thermogravimetric analyzer (DSC/TGA). The samples were heated at 5, 10 and 15 °C/min from room temperature to 700 °C in an environment comprised of 20% O₂/80% Ar by volume. Approximately 15 mg samples were loaded into crucibles and placed into the STA. Sintering can occur during heating and melting, ultimately affecting heat transfer in the STA measuring head. To ensure consistency, repeatability and minimize artifact alterations of results, four experiments were performed for each sample. Temperature calibrations for the instrument were performed using melting of a set of metal standards resulting in a temperature accuracy of ±1°C.

During the STA thermal cycle, gas species were identified using a Netzsch Aeolos 403 C quadruple mass spectrometer in order to probe for the onset and relative magnitude of hydrogen fluoride (HF) gas. Probes were programmed to collect the following masses: HF (19), H₂O (18), CO₂ (44), CF₂ (50) and CF₃ (69) but only significant values of HF were detected. Each spectrum is loaded into the analysis program as a gas signal as a function of temperature where onset and magnitude can be identified. Adiabatic flame temperature simulations for all samples were calculated using REAL code (Timtec L.L.C.). Using this thermal equilibrium software model, adiabatic flame temperature (T_f) was calculated assuming a constant pressure of 101.325 MPa and internal energy equal to zero kJ/kg. These T_f values are an upper limit for the ideal case of complete combustion. Kappagantula et al. used this software to calculate adiabatic flame temperature and their calculated values showed good agreement with their measured values for T_f (K. Kappagantula, Crane, and Pantoya 2013; K. Kappagantula, Crane, and Pantoya 2014).

Results

Flame speed results are shown in Fig. 10 and Table 2. The results show an increase of 48% from 80 nm to 120 nm and a decrease of 86% for the micron-scale AI particle size.



Figure 10. Flame speeds and activation energies for AI-PFPE as a function of AI particle size, equivalence ratio held constant and environment is air at standard temperature and pressure.

Al diameter	Adiabatic flame temperature	Peak Temperature at 5 KPM	Peak Temperature at 10 KPM	Peak Temperature at 15 KPM	Activation Energy	Flame Speed
(nm)	(К)	(°C)	(°C)	(°C)	(kJ/g)	(cm/s)
80	3900	767.7	670.0	593.8	1.53	30.5
100	3954	816.4	672.6	593.2	1.42	35.3
120	3992	819.7	677.1	589.2	1.39	45.0
5500	4063	810.0	673.6	636.3	1.72	3.3

Table 2. Adiabatic flame temperature, exothermic peak temperatures for differentDSC heating rates and calculated activation energy for all AI-PFPE samples.

Figure 11 shows the heat flux curves as a function of temperature for all particle

sizes.



Figure 11. DSC curves of the (a) pre-ignition reaction and (b) main reaction for Al-PFPE blends, Al particle diameter=80,100,120,5500 nm in 20% O₂/80% Ar by volume. Heating rate is 15 KPM for all data presented. Data reporting specific energy for each exotherm.

The exotherms for the PIR and main reaction increase as particle size

increases from 80 to 120 nm and both exotherms decrease for the micron scale

samples (see Table 3 for values).

Al diameter	PIR	PIR onset	main reaction	main reaction onset	mass loss	F gas signal per mg PFPE	Gas Signal Onset
(nm)	(J/g)	(°C)	(J/g)	(°C)	(%)	(A*s/mg)	(°C)
80	19.6	315.6	123.5	558.5	-68.9	27.1	351.0
100	19.8	314.7	132.5	560.8	-68.8	26.7	341.7
120	21.0	312.9	146.2	564.0	-68.8	23.9	349.9
5500	7.3	313.4	73.0	618.2	-57.2	20.0	363.1

Table 3. Magnitude and onset of exotherms, gas curves and mass loss for each sample.

The onset for the micron-scale main reaction is about 60 °C higher than the main reactions for the nano-scale reactions. These results show that micronscale AI is less reactive than nano-scale counterparts and this has been seen by others(Levitas, Dikici, and Pantoya 2011; Hunt et al. 2009; M. Pantoya and Granier 2005; Hunt and Pantoya 2005).

Figure 12a shows gas species evolution for HF liberation in each sample. It is noted the signals for H_2O , CO_2 , CF_2 and CF_3 only account for 2% of the total gas signal. The area under each curve (Fig. 12a) was normalized to the mass of PFPE in the systems and that data is shown in Fig. 12b. The normalized gas signal decreases as particle size increases (i.e., as the alumina concentration decreases).



Figure 12. (a) HF gas evolution for 80, 100, 120 and 5500 nm Al with PFPE and (b) gas signal per mg of PFPE in each sample. The amount of HF liberated from each sample decreases as particle size increases (i.e., as alumina content decreases).

Table 2 shows the adiabatic flame temperatures calculated via thermal-

equilibrium software. As particle size increases from 80 to 5500 nm, the

adiabatic flame temperature increases from 3900 to 4016 K.

Activation energy is calculated from the DSC curves using the

isoconversion method in Eq. (11)(K. S. Kappagantula et al. 2012; Hunt and

Pantoya 2005; Farley et al. 2013), where B is the DSC heating rate (5, 10, and 15

Kelvin per minute (KPM)) and T_{ρ} is the temperature of the peak of the exothermic reaction (given in Table 2).

$$\ln\left[\frac{B}{T_p^{1.95}}\right] = -\frac{E_a}{RT_p} + lnA \qquad (11)$$

By measuring T_p for three heating rates, E_a can be extrapolated from the slope for a mixture (see Fig. 10).



Figure 13. Trend lines showing activation energy for each particle diameter Al-PFPE samples. Activation energies are reported in Table 2.

This and similar proposed methods for calculating activation energy have

been used by others(Kissinger 1957; Ozawa 1992; Boswell 1980; Starink 1996;

Dyer and Munir 1995).

Discussion

Figure 10 shows that the fluorinated coating is significantly more reactive

with nano-scale Al particles than micron scale. This may be associated with the

exposed Al_2O_3 surface area of each sample. The Al particles are spherical and the specific surface area (1/nm) can be calculated by Eq. (12).

$$A_{sp} = \frac{4\pi r^2}{\frac{4}{3}\pi r^3} = \frac{3}{r}$$
(12)

The magnitude of the PIR increases in nano-AI-PFPE as specific surface area decreases (Tables 1 and 3). Also, the magnitude of the PIR correlates with flame speed (Fig. 10 and 11a): larger nano-AI particles produce faster flame speeds with PFPE. This correlation implies that optimizing the calorific output from the PIR can lead to enhanced energy propagation. Once the average particle size reaches the micron-scale (i.e., 5500 nm AI), flame speeds decrease, in part, due to insufficient alumina present to catalyze PFPE decomposition.

The flame speed trends (Fig. 10) also follow closely with the main reaction exotherm magnitude, specifically; higher flame speeds correspond to larger exotherms. These results (Fig. 11b) suggest a correlation between calorific output and energy propagation, implying a dominant role of conductive energy propagation. These blends are highly consolidated mixtures with bulk density ranging from 87 to 92 %TMD such that porosity is limited to 13-8% of the solid volume. For these high density blends convective energy transport is more limited and conduction is dominant. The samples with larger magnitude PIR and

main reactions may provide more energy for conductive heat transfer thereby exhibiting high flame speeds.

Activation energies for all AI-PFPE blends correlate well with the inverse of the measured flame speeds (see Table 2). As activation energy decreases, flame speed increases suggesting flame speed is a function of the reaction kinetics as well as the conductive mode of energy propagation.

Because the PIR is highly controlled by the specific surface area, a way to investigate the extent of the surface chemistry is to vary the aluminum particle size, and therefore exposed surface area. As the specific surface area increases (i.e. particle size decreases), the available sites for AI-F surface reactivity increase. Kasai et al. showed that alumina is a good catalyst for PFPE decomposition¹⁴. Because the mixtures were blended to a constant stoichiometry, the alumina concentration increases as particle size decreases. This is why decreasing AI particle size produces an increase in specific HF gas production (see Fig. 12). The PFPE is exposed to more alumina surface area and undergoes better decomposition with the 80 nm particle size. Because the equilibrium analysis and open tray burns allow for much of the HF gas to escape the system, much of the fluorine that would catalyze surface combustion has left the system prior to energy propagation. The decrease in performance in the micron-AI-PFPE may be

caused, in part, by the higher activation energy allowing appreciable HF gas escape prior to particle ignition and energy propagation.

Conclusion

Flame speeds, differential scanning calorimetry (DSC), thermogravimetric analysis (TGA) and quadruple mass spectrometry (QMS) were performed for Al-PFPE blends with varying Al particle sizes (80, 100, 120 and 5500 nm in average diameter). The results show that the combustion performance of these blends is highly dependent on the Al_2O_3 exposed surface area. High specific surface area associated with nano-AI particles catalyzes AIF₃ formation via exothermic surface chemistry. This reaction promotes the decomposition of PFPE (as seen by higher HF gas evolution), and provides greater overall calorific output for diffusive energy propagation, as seen in the higher flame speeds for nano-Al-PFPE blends. Activation energy was also measured and found to inversely correlate with flame speed indicating that reaction kinetics also play a dominant role in improving combustion performance. These results provide a new direction for activating Al particles towards greater reactivity: by including a fluorinated oligomer to exploit exothermic surface reactions contributing toward overall energy generation.

<u>CHAPTER III: Activating Aluminum Reactivity for Improved Energetic Composite</u> <u>Combustion</u>

Abstract

Energetic blends of nanometer-sized aluminum (AI) particles with liquid perfluorocarbon-based oxidizers such as perfluoropolyethers (PFPE) excite surface exothermic chemistry between fluorine and the alumina passivation shell surrounding an Al core particle that promotes Al reactivity. Many Al fueled energetic composites use solid oxidizers that induce no alumina surface exothermicity, such as molybdenum trioxide (MoO3) or copper oxide (CuO). This study investigates a perfluorinated polymer additive, PFPE, incorporated to activate AI reactivity in AI-CuO and AI-MoO3. Flame speeds, differential scanning calorimetry (DSC) and quadruple mass spectrometry (QMS) were performed for varying percent PFPE blended with Al/MoO3 or Al/CuO in order to examine reaction kinetics and combustion performance. X-ray photoelectron spectroscopy (XPS) was performed to identify product species. Results show that the performance of the thermite-PFPE blends is highly dependent on the bond dissociation energy of the metal oxide. Fluorine-aluminum based surface exothermic chemistry with MoO3 produce an increase in reactivity while the blends with CuO show a decline when increasing the PFPE loadings. These results

provide new evidence that optimizing aluminum combustion can be achieved through activating exothermic Al surface chemistry that produces aluminum fluoride.

Introduction

A thermite is defined as an energetic system consisting of a metal fuel (i.e. aluminum (AI)) and metal oxide (i.e. copper oxide (CuO) or molybdenum trioxide (MoO₃))(Nixon et al. 2010; Gesner, Pantoya, and Levitas 2012; Fischer and Grubelich 1998; Hunt and Pantoya 2005). These systems are widely studied due to their high energy densities and heats of combustion(Cervantes et al. 2011; Yen and Wang 2012; Kyle T. Sullivan et al. 2010; M. Pantoya and Granier 2005; Hunt and Pantoya 2005; Sanders et al. 2014). As energy generating materials, thermite reactions have applications ranging from material synthesis(Shigeta and Watanabe 2010; Deevi, Sikka, and Swindeman 1997; Yeh and Huang 2011) and alloying(Charles a. Crane, Pantoya, and Dunn 2010; L. L. Wang, Munir, and Maximov 1993) to welding and joining(lordachescu et al. 2010; Chen et al. 2006). Munir et al. present a thorough review on thermite applications(Dyer and Munir 1995), Fischer and Grublic tabulated thermo-chemical properties for hundreds of thermite reactions (Fischer and Grubelich 1998) and Koch presented chemistry and application of fluorocarbon-based energetics(Koch 2012).

For Al-based thermites, when fluorine replaces oxygen as the reducing agent, aluminum fluoride (AIF₃) is formed instead of Al₂O₃. Both AlF₃ and Al₂O₃ are comparably thermodynamically stable with heats of formation of 1510 kJ/mol and 1676 kJ/mol, respectively. However, AlF₃ formation is preferred to Al₂O₃, because Al-F (664±6 kJ/mol)(Cottrell 1958) bond formation is stronger than the Al-O bond (512±4 kJ/mol)(Cottrell 1958). In fact, F has been shown to react with the alumina passivation shell surrounding an Al particle (Osborne and Pantoya 2007). The exothermic surface reaction precedes the main Al oxidation reaction and has been coined a pre-ignition reaction (PIR)(Osborne and Pantoya 2007). Kappagantula et al. showed that the PIR can be used to enhance Al reactivity(K. S. Kappagantula et al. 2012).

Typically Al-fluoropolymer blends are prepared with a solid fluorinated oxidizer such as Teflon[®] (PTFE) (K. Kappagantula, Pantoya, and Hunt 2012; M. L. Pantoya and Dean 2009; Losada and Chaudhuri 2010; Conner and Dlott 2012). Solid PTFE introduces a limit to the fuel-oxidizer contact surface area which is thought to limit the combustion performance(K. S. Kappagantula et al. 2012). In an attempt to resolve the limitation caused by fuel-oxidizer contact surface area, Kappagantula et al(K. S. Kappagantula et al. 2012) investigated the combustion of Al particles with surface functionalized self-assembled monolayers (SAM). The SAM consisted of perfluorotetradecanoic (PFTD) acid bonded to the aluminum particle's native oxide shell (Al₂O₃). Surface functionalized Al-PTFD was successful in increasing the reactivity of Al particles combined with MoO₃ and produced an 86% increase in flame speed compared to Al-MoO₃ alone(K. S. Kappagantula et al. 2012). The enhancement in reactivity was attributed to the PIR resulting from the fluorine functionalization.

Another method for coating the Al surface that can be easily implemented is to introduce a liquid perflurorinated oligomer that phyisoabsorbs onto the Al particle surface. In a similar finding as Kappagantula et al., the liquid fluorooligomer may readily activate the PIR and promote greater overall Al reactivity. The liquid fluorooligomer used in this study is a class of viscous fluorinated oligomers called perfluoropolyethers (PFPEs). Miller et al. used a PFPE protective coating on nano-scale Al because of its thermal stability at low temperatures (up to 316 °C in an oxygen-rich environment) and potential to produce exothermic activity beyond 316 °C(Miller et al. 2013). When coated with PFPE, the Al particles were able to suspend in a structural epoxy-based matrix. Miller used differential scanning calorimetry (DSC) to study the exothermic behavior for Al-perfluoropolyether (PFPE) blends of varying weight percents. The DSC revealed two exothermic peaks, one of which occurred at the PFPE

decomposition temperature of 316 °C (i.e., the PIR). The 30% AI 70% PFPE blend produced the highest heat of combustion. This translated to an equivalence ratio (ER) of 1.2 (i.e., slightly fuel rich).

Previously, fundamental combustion analysis was performed for AI-PFPE blends(McCollum, Pantoya, and Iacono 2015). Flame speeds, differential scanning calorimetry (DSC), thermogravimetric analysis (TGA) and quadruple mass spectrometry (QMS) were performed for AI-PFPE blends with varying AI particle sizes (80, 100, 120 and 5500 nm in average diameter). The results showed that the combustion performance of these blends is highly dependent on the Al₂O₃ concentration and exposed surface area. As Al particle diameter increases from 80 to 120 nm, flame speeds increase by 48%, but from 120 to 5500 nm, flame speeds decrease by 93%. There is a balance to optimizing Al particle reactivity between activating Al particles with exothermic surface chemistry versus the unreacted alumina that contributes a thermal heat sink during energy generation. These results introduced a new and simplified synthesis approach for activating Al particle surface chemistry to promote overall reactivity.

The objective of this study is to use PFPE as an additive in different thermite systems (Al-MoO₃ and Al-CuO) in order to determine how the

perfluorinated oligomer affects reaction kinetics and overall reactivity. The metal oxides MoO₃ and CuO were selected based on their extensive use in research (Jian, Piekiel, and Zachariah 2012; H. Wang et al. 2014; K. Sullivan, Young, and Zachariah 2009; D. S. Moore, Son, and Asay 2004; Sanders et al. 2014) and their differing dissociation properties. Specifically, the dissociation energy for the Mo-O bond is 607±34 kJ/mol(Cottrell 1958) but the Cu-O bond energy is only 343±63 kJ/mol(Cottrell 1958). Using a PFPE coating in these thermite systems could extend their shelf life by limiting particle surface exposure to oxygen rich environments while exploiting exothermic surface chemistry with the alumina shell that may contribute to enhancing overall reactivity.

Experimental

Materials

Aluminum powder (100 nm average particle diameter) with 3 nm Al₂O₃ particle shell thickness was procured from U.S. Nanomaterials (Houston, TX). The PFPE (Fomblin® Y LVAC 25/6, average molecular weight of 3300 g mol⁻¹) and the CuO (50 nm average particle diameter (spherical)) powder was purchased from Sigma Aldrich (St. Louis, MO). The MoO₃ (44 nm average particle diameter (flake structures)) was received from Nanostructured & Amorphous Materials, Inc. The composites were prepared to an equivalence ratio of 1.2 (i.e., slightly fuel rich). Equivalence ratio is shown in Eq. (13) where M is the mass and the *fuel* is Al, and the *oxidizers* are PFPE/CuO or PFPE/MoO₃.

$$\mathbf{ER} = \frac{\left[\frac{M_{fuel}}{M_{oxidizer}}\right]_{actual}}{\left[\frac{M_{fuel}}{M_{oxidizer}}\right]_{stoichiometric}}$$
(13)

The equivalence ratio was maintained for all samples by performing a molar balance using Eq. (14) and Eq. (15) where a, b, c, d and e are coefficients used to balance each reaction and H represents other products of the reaction.

Or,

aCF₃O-[-CF(CF₃)CF₂O-]_x(-CF₂O-)yCF₃ + bAl + cCuO
$$\rightarrow$$

dAl₂O₃ + eAlF₃ + H (15)

The PFPE concentration was calculated such that PFPE accounts for 0, 5, 10 and 20 wt. % of the reaction. The weight percent of each component are shown in Table 4.

	Al	Oxidizer	PFPE
Al-CuO		CuO	
0	24%	76%	0%
5	23%	73%	5%
10	22%	69%	9%
20	21%	63%	16%
Al-MoO ₃		MoO ₃	
0	34%	66%	0%
5	34%	62%	4%
10	33%	59%	8%
20	32%	51%	17%

Table 4. Weight percent of all components in each sample.

The PFPE, AI and CuO or MoO₃ were weighed and suspended in 60 ml of Perfluorosolv PFS-2[™] from SPI Supplies (West Chester, PA). A fluorinated solvent for mixing was included in order to improve PFPE dispersion. The fluorine in the solvent disperses the fluorine in the PFPE and suspends the AI/CuO or AI/MoO₃ particles to promote better coating. The solution was then mixed using a planetary mixer at 1500 rpm for 2 minutes and poured into a Pyrex dish. The Perfluorosolv evaporated in a fume hood until the remaining mass was only that of AI/CuO/PFPE or AI/MoO₃/PFPE (about 24 h). Because of the high surface area of the AI/CuO and AI/MoO₃ and low concentration of PFPE, the samples reclaimed as loose powders.

Flame Speed Measurements

The powder was loaded into 3 mm inner diameter, 8 mm outer diameter, 10 cm long quartz tubes containing 300 mg of powder each (Fig. 14).



Figure 14. Representative still frame images time stamped of powder filled quartz tube and flame propagation. Bulk density is 10% theoretical maximum density and mixture is Al/MoO₃.

Both ends of the tube were sealed with one side securing a length of

nickel-chromium wire, bent into a "V" shape and used for ignition by subjecting

the wire to a voltage. Flame propagation was observed through a viewing

window in the chamber. The reaction was recorded with a Phantom v7 (Vision

Research, Wayne, NJ) high speed camera at a rate of 29,000 frames per second

and aligned perpendicular to the direction of flame propagation. The flame speed was determined by tracking the flame front through a referenced time and distance using the Vision Research Software. The resolution of the flame speed for this diagnostic is 0.1 m/s. Flame speeds were collected from five sets of experiments to ensure repeatability.

Thermal Equilibrium Analysis

Simultaneous Thermal Analysis (STA) was performed using a Netzsch Jupiter STA 449 differential scanning calorimeter and thermogravimetric analyzer (DSC/TGA). The samples were heated at 5 °C/min from room temperature to 1000 °C. Approximately 10 mg samples were loaded into crucibles and placed into the STA. Sintering can occur during heating and melting, ultimately affecting heat transfer in the STA measuring head. To ensure consistency, repeatability and minimize artifact alterations of results, five experiments were performed for each sample. Temperature calibrations for the instrument were performed using melting of a set of metal standards resulting in a temperature accuracy of ±1°C.

Gas species were identified using a Netzsch Aeolos 403 C quadruple mass spectrometer in order to probe for the onset and relative magnitude of gaseous fluorine during the STA thermal cycle. Each spectrum is loaded into Netzsch Proteus software in order to identify species and magnitude as a function of temperature. Probes were programmed to collect the following masses: HF (19), H_2O (18), CO_2 (44), CF_2 (50) and CF_3 (69). Because H_2O , CO_2 , CF_2 and CF_3 only made up 2% of the total gas signal, the focus will be on HF.

X-ray Photoelectron Spectroscopy

X-ray photoelectron spectroscopy (XPS) was performed on the reaction products using a PHI 5000 Versa Probe with an Al Kα source. Samples were loaded in a vacuum chamber which was held to 10⁻⁶ Pa during measurement. Peaks were referenced to a C1s value of 284.8 eV. For the MoO₃-based thermite (MBT) samples, the survey spectra identified Al, Mo, O, F, and C. The CuO-based thermite (CBT) survey spectra identified Al, Cu, O, F, C and Na. Due to the presence of Na, analysis was performed on the O2s peak because the O1s peak was impeded by a Na1s peak.

Results and Discussion

Flame speed results for the two thermite systems as a function of PFPE concentration are shown in Fig. 15.



Figure 15. Flame propagation velocity for aluminum with (a) CuO and (b) MoO₃ with varying percent of PFPE. Equivalence ratio remains constant for all mixtures.

Addition of PFPE yields different results with regard to oxidizing agent. The MoO₃-based thermites (MBT) benefit from the addition of PFPE, showing a continuous increase in flame speed (37% with the highest PFPE concentration). The CuO-based thermites (CBT) show the opposite trend: flame speed decreases as the PFPE concentration increases, with a decrease of 7% for the highest PFPE concentration.

The key question in this analysis is how fluorine from PFPE participates in the reactions. When combined with aluminum, the resulting product will be Al_2O_3 if AI reacts with oxygen reducing agents (i.e., CuO or MoO₃) and AIF₃ if AI reacts with the PFPE. The equivalence ratio (Φ =1.2) was maintained with the addition of PFPE such that the oxygen and fluorine have equal availability to react with Al. Figure 16 shows the thermal analysis for 100 nm Al and PFPE alone. The results for onset and heat of combustion of each peak are shown in Table 5.



Figure 16. The oxidation reaction of 100 nm Al and PFPE heated from 35 to 1000°C at 5°C/min.

Table 5. Results for the DSC analysis of Al-CuO and Al-MoO₃ with x% of PFPE.

Sample	PIR Onset	PIR ΔH _c	Thermite Reaction	Thermite Reaction
	Temperature (°C)	(J/g)	Onset Temperature	ΔH _c
		-	(°C)	(J/g)
AI + PFPE	315	19.8	561	133
AI + CuO +0%PFPE			517	763
AI + CuO +5%PFPE	298	19.56	569	1658
AI + CuO +10%PFPE	299	29.70	581	1305
AI + CuO +20%PFPE	303	51.37	583	843
AI + MoO ₃ +0%PFPE			508	2078
AI + MoO ₃ +5%PFPE	298	21.79	534	1370
AI + MoO ₃ +10%PFPE	301	35.17	541	1672
AI + MoO ₃ +20%PFPE	305	103.1	566	1889

Figure 17 shows heat flow curves from thermal analysis for CBT and MBT from 280-380 °C. The magnitude of the CBT pre-ignition reaction (PIR) builds with PFPE loading (Fig. 17a), but is significantly less than that of the MBT (Fig. 17b). Also, the onset of the PIR is around 307 °C for both CBT and MBT. Table 5 shows the magnitudes of the PIR for each sample.



Figure 17. Pre-ignition reaction exotherm for aluminum with (a) CuO (PIR region) or (b) MoO_3 (PIR region) and (c) MBT and CBT (400-700 °C region). Samples were heated at 5 °C/min from 35°C to 1000°C in a 20% $O_2/80\%$ Ar by volume environment.

This low temperature exotherm has been identified by others and is associated with AIF_3 formation reactions(K. S. Kappagantula et al. 2012; Travis R. Sippel, Son, and Groven 2013; Miller et al. 2013; Farley et al. 2013).

The thermal analysis from 400-700°C for all samples is shown in Fig. 17c and the results are summarized in Table 5. The onset temperature of the main reaction increases with both CBT and MBT as PFPE concentration increases. Also, the magnitude of the main reaction produces a continuously increasing heat of combustion (calculated as the area under the curve) with increasing PFPE for both metal oxide blends. As the amount of PFPE increases, both onset temperatures surpass the onset temperature for Al oxidation at 565 °C when heated with PFPE alone (Fig. 16). It is noted that PIR are not shown in Fig. 17c because the difference in magnitude between the PIR and main reaction is too large to distinguish the smaller PIR peaks.

From Figs. 15-17, an increase in flame speed was observed in for MBT, and MBT shows that the PIR grows in exothermicity as PFPE concentration increases (Fig. 15b). This suggests that fluorine contribution to the reaction increases as the PFPE concentration increases and this growing PIR may correlate to an increase in flame speed with PFPE concentration. In contrast, Figs. 15-17 shows CBT exhibits a decrease in flame speed with PFPE concentration and the PIR is

significantly smaller than that of the MBT samples (see Table 5). Without significant contribution from the early stage PIR (Table 5), chemical energy liberated from the reaction is delayed to higher onset temperature thereby limiting energy propagation and reducing flame speed.

The opposite trends in flame speed shown in Fig. 15 may depend on the bond dissociation energy (BDE) of the different metal oxidizers. The energy to break the Cu-O bond is only 343±63 kJ/mol but for the Mo-O bond it is 607±34 kJ/mol(Cottrell 1958). The fluorine received from the PFPE is bonded to carbon and the BDE of that bond is 536±21 kJ/mol(Cottrell 1958), the carbon-carbon bond is 607±21 kJ/mol(Cottrell 1958) and the carbon-oxygen bond is 1076.5±4 kJ/mol(Cottrell 1958). Crouse et al. suggested that a possible mechanism for Al₂O₃ catalyzed reaction with PTFE involves the chemisorption of a CF₂ radical(Puts and Crouse 2014). The energy needed to dislodge an oxygen atom from CuO to react with Al is significantly lower than the energy needed to extract fluorine alone from PFPE or to extract the CF₂ radical by breaking the C-C bond in the PFPE. For this reason, the PFPE reaction with AI may be overshadowed in the CBT because O is more readily available to oxidize Al (i.e., 343 vs. 607 m/s). However, when the blends are made with an oxidizer with a higher BDE (i.e. MoO_3), the PFPE has a greater chance to participate in the reaction because the

energy needed to extract fluorine (or CF_2 radical) and oxygen atoms from their respective reducing agents are similar. This can be confirmed by analyzing the products of the different reactions in order to identify the presence of AIF_3 in the different samples.

Hydrogen fluoride (HF) gas generation was monitored during thermal analysis and shown as a function of temperature in Fig. 18a and 18b. For CBT, HF gas evolution decreases with increasing concentration of PFPE. In contrast, MBT show greater HF gas evolution with increasing PFPE concentrations. Figure 17c shows the HF gas signal per mg PFPE (As/mg) for all samples.



Figure 18. QMS curves for fluorine gas liberated from (a) CBT, (b) MBT and (c) gas evolution per mg PFPE. The MBT show an increase and limit in HF gas liberation while the CBT show a decrease.

The onset of HF gas release is approximately < 350 °C which corresponds to the decomposition of PFPE(Kasai, Tang, and Wheeler 1991). The CBT show a decrease in the release of HF per mg of PFPE which indicates that F bonds are either being formed with the other materials in the system, or that PFPE decomposition is hindered in some way. The normalized spectrum for the MBT show an increase in HF gas produced per mg PFPE from 5 to 10%, but there is little change from 10 to 20% PFPE concentration. This data suggests that more Al-F bonds are formed with the MBT than the CBT and is consistent with the magnitudes of the PIR in Fig. 17 and Table 5. Specifically, more exothermic PIR is consistent with more Al-F bond formation and lower HF gas evolution.

In order to identify how F bonds to each component in the combustion products, XPS was performed on products from the MBT and CBT reactions for 5% and 20% PFPE. Figure 19 shows the survey spectrum of the MBT samples (i.e., a. MBT5, b. MBT20) and CBT samples (i.e., c. CBT5 and d. CBT20).



Figure 19. Survey Spectra for (a) MBT5 and (b) MBT20 (c) CBT5 and (d) CBT20.
The relative signal of fluorine decreases significantly from CBT5 to CBT20. (Note: The counts per second axis of the CBT samples are shifted in order to see the low intensity peaks in CBT20.)

Fluorine bonds are present for all samples. Inspection of these plots shows a comparable signal of every element in the MBT samples, but a drastic decline in F signal from CBT5 to CBT20. Upon closer inspection, Mo3d shows a greater number of pronounced peaks in the MBT5 sample versus MBT20. The peaks at 226 eV and 229 eV are metallic Mo 3d5/2 and 3d3/2 respectively (see Fig. 20)(Al-Kandari et al. 2013). Peaks at 228, 231, 232, and 235 are indicative of Mo-O bonds(Al-Kandari et al. 2013).



Figure 20. Mo3d spectral line for (a) MBT5 and (b) MBT20.

On the other hand, the CBT show only metallic Cu2p peaks at 232 and 252 eV for the 3/2 and 1/2 spins respectively (see Fig. 21)(Espinós et al. 2002). (Note: There is a small peak at 945 eV in both Cu spectra. This is a Cu3p3/2 satellite peak (Parmigiani et al. 1992).) This also shows that there is no bonding between Cu and F.



Figure 21. Cu2p spectral line for (a) CBT5 and (b) CBT20.

Also, due to the presence of the Cu3s peaks, tight scans of the strong Al2p peak were not used because they fall in the same binding energy range, (Herrera-Fierro 1993) instead Al2s was analyzed. There is still a Cu3s peak in the same region, but because there is only one peak from Cu, and the binding energies are spaced well enough to see 2 distinct peaks, Al2s was the better candidate for peak identification. For CBT samples, the Al2s peak was comprised of Al-O (119 eV) and Al-F (121 eV) binding energies.(see Fig. 22c and 22d)(Hess et al. 1993). It is noted that the peak located at 122 eV in the CBT5 and CBT20 Al spectra is the Cu3s peak(Parmigiani et al. 1992). The Al2s peak was used for both CBT samples

and Al2p was used for the MBT samples. The Al2p line showed Al-O bonds at ~74.0 eV and Al-F at 75.2 eV (Fig. 22a and 22b). Figure 22e shows the ratio of Al-F peak intensity to Al-O peak intensity for each sample.





(e)

Figure 22. Al2p spectra for (a) MBT5 and (b) MBT20 and Al2s spectra for (c) CBT5 and (d) CBT20 and (e) graphical representation of Al-F bonds to Al-O bonds present in each sample.

Although Al-F bonds are detected in all samples, the MBT samples show an increase in the ratio of Al-F/Al-O bonding while the CBT samples show a decrease. The increase in Al-F bonding was expected with increasing PFPE loading, but the CBT samples showed a decline. This decline along with the decreasing HF gas signal from CBT5 to CBT20 provides new information about the location of the fluorine.

In addition to AI-F bonds present in CBT samples the F1s line shows an additional peak at 688 eV(Herrera-Fierro 1993) for CBT5 and 690 eV(J. M. Lee et al. 2009) in the CBT20 sample (see Fig. 23c and 23d). These peaks are both indicative of C-F bonds and were only distinguished in the CBT samples. The MBT samples only show one fluorine peak at 686 eV (Fig. 23a and 23b).



Figure 23. F1s spectra for (a) MBT5, (b) MBT20, (c) CBT5 and (d) CBT20.

This along with the increase in HF gas production suggests that the Al-MoO₃ blends catalyze the decomposition of PFPE while the Al-CuO blends do not. This is seen in the F1s spectra for CBT versus MBT. The improved decomposition of PFPE in the MBT samples along with the increasing formation of Al-F bonds confirm that the low temperature surface chemistry shown by the PIR increases flame speed in these samples.

Conclusion

Combustion characterization analyses using high speed imaging, XPS, QMS, and DSC were performed for varying concentrations of PFPE in Al/MoO₃ and Al/CuO composites in order to examine the influence of fluorine surface chemistry on reactivity. Results show that the performance of the thermite-PFPE blends is highly dependent on the oxidizing agent. The PFPE blends with MoO₃ show an increase in reactivity while the blends with CuO show a decrease in reactivity when increasing PFPE concentration. We observed a decline in formation of AlF₃ in CBT samples but an increase in AlF₃ formation in the MBT samples. The potential surface chemistry between Al₂O₃ and F from PFPE is negated by competitive Al-O, but thermodynamically favorable formation from CuO with lower bond dissociation energy than F or the CF₂ radical. When using an oxidizer with a BDE similar to that of the C-C or C-F bond (i.e. MoO₃), PFPE can

promote reactivity via catalytic behavior of the alumina shell to help decompose PFPE more efficiently and improve the low temperature surface chemistry and overall reactivity.

CHAPTER IV: Improving Aluminum Particle Reactivity by Annealing and Quenching Treatments: Synchrotron X-ray Diffraction Analysis of Strain

Abstract

In bulk material processing, annealing and quenching metals such as aluminum (AI) can relieve residual stress and improve mechanical properties. On a single particle level, affecting mechanical properties may also affect AI particle reactivity. This study examines the effect of annealing and quenching on the strain of AI particles and the corresponding reactivity of aluminum and copper oxide (CuO) composites. Micron-sized AI particles were annealed and quenched according to treatments designed to affect AI mechanical properties. Synchrotron X-ray diffraction (XRD) analysis of the particles reveals the thermal treatment increased the dilatational strain of the aluminum-core, alumina-shell particles. Flame propagation experiments also show thermal treatments effect reactivity when combined with CuO. An effective annealing/quenching treatment for increasing aluminum reactivity was identified. These results show that altering the mechanical properties of AI particles affects their reactivity.

Introduction

Composite energetic materials consist of a metallic fuel (i.e., aluminum (AI)) and a metal oxide (i.e., copper oxide (CuO)). These composites are widely studied due to their high energy densities and heats of combustion. A main goal of much published research studying aluminum combustion is to understand and improve particle reactivity. Towards this end, many have proposed new Al synthesis strategies: such as altering the native aluminum oxide (Al2O3) coating with another passivating agent such as alkenes (Chung et al. 2009) or applying self-assembled monolayers (SAM) to the particle surface(K. S. Kappagantula, Pantoya, and Horn 2013). Various explanations for Al oxidation mechanisms have also been proposed, each strongly tied to the ignition mechanism and heating rate(Levitas et al. 2006; Levitas et al. 2007; M. A. Trunov, Schooenitz, and Dreizin 2006; Dreizin 1999; M. a Trunov et al. 2006; K.T. Sullivan et al. 2012; Rai et al. 2006; K. Sullivan and Zachariah 2010; Chakraborty and Zachariah 2014). All theories share a common theme for mass transport of fuel and oxidizer, but differ in how that diffusion is achieved (i.e., via (a) dispersion(Levitas et al. 2006; Levitas et al. 2007), (b) phase changes in the polymorphous passivation shell(M. A. Trunov, Schooenitz, and Dreizin 2006; Dreizin 1999; M. a Trunov et al. 2006), (c) reactive sintering(K.T. Sullivan et al. 2012), (d) pressure gradient driven

processes(Rai et al. 2006; K. Sullivan and Zachariah 2010), and (e) induced electric field influences(Chakraborty and Zachariah 2014)). This article will not directly deal with any particular reaction mechanism, but rather investigate the influence of a new parameter, mechanical strain, which has only recently been considered in the study of Al oxidation(Levitas et al. 2006; Levitas et al. 2007; Levitas, Mccollum, and Pantoya 2015; Dikici, Pantoya, and Levitas 2010).



Figure 24. Transmission electron spectroscopy (TEM) image of a nano-scale Al particle to show alumina shell thickness. The average thickness is 4.01 nm. This thickness was used in the purity calculations for micron-scale Al particles (< 1% by mass oxide).

A typical aluminum particle consists of an aluminum core passivated by an

alumina shell as seen in Fig. 24. Residual stress within an aluminum particle is

induced during production. These are thermally induced stresses that are

generated during the cooling stage. Stress builds in two stages during particle

synthesis: (1) within the aluminum core during cooling and prior to oxide shell formation; and, (2) during alumina shell formation until the core-shell system reaches ambient temperature conditions.

Within the molten aluminum droplet absent of a shell, stress develops as the molten aluminum droplet cools, because external surface layers start to shrink while the core is still hot and free to contract but its contraction is constrained by the external layers that have solidified and become rigid. In this way, the stress distribution in a naked aluminum particle is tensile in the core and compressive at the surface. Smaller particles cool at faster rates(Zheng et al. 2009a; Zheng et al. 2009b; Ozbilen, Unal, and Sheppard 2000) and should manifest larger internal stresses during production.

The shell is formed when oxygen is introduced at temperatures below the melting temperature of aluminum (i.e., < 660 °C(Hatch 1984)) and no higher than 440 °C (i.e., the phase transition of amorphous to gamma alumina(Dorre and Hubner 2011)). At this point, another residual stress develops because there exists a mismatch of thermal expansion coefficients (i.e., a, linear coefficient; or, b, volumetric coefficient) between the newly formed alumina shell (i.e., $a_{ox} = 5 \times 10^{-6} \text{ K}^{-1}$ and box= 8 x 10⁻⁶ K⁻¹ (Dorre and Hubner 2011)) and aluminum core ($a_m = 23 \times 10^{-6} \text{ K}^{-1}$ and $b_m = 69 \times 10^{-6} \text{ K}^{-1}$ (Hatch 1984)). Cooling from 440 °C to ambient

will induce shrinkage in the core while the shell remains relatively rigid, i.e., there is an order of magnitude difference in expansion coefficients. The internal radius of the oxide sphere is forced to decrease and stresses arise at the interface of the core-shell.

Isothermal stress relief is a common approach to stress relaxation(Throop, Underwood, and Legar 1971; James 1987; L. W. Crane 1979). This technique involves the uniform heating of a material to a specified temperature below the melting temperature, holding at that temperature for a period of time, followed by cooling to control the re-introduction of desirable thermal stresses. This sequence of steps is referred to as annealing followed by quenching. Timoshenko and others(Throop, Underwood, and Legar 1971; James 1987; L. W. Crane 1979; Vilms and Kerps 1982; Kingery 1955) explain significant microstructural changes take place that promote stress relief at about two-thirds the temperature at which the stresses were formed. In the case of Al particles, the amorphous alumina shell is formed at roughly 440 °C such that annealing temperatures should at least be 293 °C. Interestingly, Firmansyah et al. examined the stress state of nano-aluminum powder upon continuous heating using a high temperature XRD and found that AI particles experience a zero-stress state at 300 °C(Firmansyah et al. 2012).

The main mechanism that causes relaxation of locked-in stresses for annealing temperatures < 400°C is classical diffusional creep. This mechanism enables counterbalancing regions of tensile and compressive stresses to contract or expand slightly, and thus to redistribute. This is also a time dependent process and determining the optimum temperature and duration for annealing and quenching has not previously been investigated for aluminum particles. Some experimental work on other metals has revealed a relevant conclusion: while the annealing time is important, creep is a logarithmic process such that most relief is obtained at a given temperature rapidly(James 1987). Data presented by Adeyemi et al. for carbon steel suggests annealing times on the order of 10 minutes should be sufficient to relieve stress(Adeyemi, Modlen, and Stark 1983). Even for the annealing temperature of 500oC, 80% of the residual stress is relieved in 5 minutes. Bulk aluminum alloys have been studied for creep behavior (Prasad, Sastry, and Vasu 1970). Prasad et al. studied creep at 87 and 200 °C and found the effects of stress increments and decrements to be different(Prasad, Sastry, and Vasu 1970). This is an important finding because it implies stress variations are significant in the temperature ranges we are most interested (i.e., < 400 °C).

Quenching is another important variable to consider. Evancho et al. showed that slow cooling may re-introduce thermal stresses that are undesirable (e.g., like the residual stress) and lead to lower strength(Evancho and Staley 1974). Faster cooling rates may effectively 'freeze' the stress state such that purposefully induced desirable thermal stress do not have time to relax, leading to higher strength. The hypothesis is: faster cooling rates will produce aluminum particles with optimal mechanical properties promoting optimal reactivity.

Dikici et al. showed preliminary evidence that annealing and quenching Al particles affects reactivity (Dikici, Pantoya, and Levitas 2010). They heated Al particles to a prescribed temperature then cooled them to room temperature and found that some thermal treatments lead to improved flame speeds. Their powders were allowed to cool at two different rates. The samples that were cooled at 0.06 KPS (Kelvin per second) showed flame speeds comparable to the untreated samples, but when the cooling rate was raised to 0.13 KPS, the flame speeds for micron scale Al and molybdenum trioxide (MoO3) improved for one case and worsened for the other. More recently, Levitas et al. extended the work by Dikici et al. by focusing on micron scale Al particles and using a different metal oxide (i.e., copper oxide (CuO))(Levitas, Mccollum, and Pantoya 2015). They examined flame speeds and showed certain thermal treatments were consistent

with predictions of the melt dispersion mechanism, effectively expanding the realm of this reaction mechanism towards optimization of micron scale Al particles (Levitas, Mccollum, and Pantoya 2015).

The objective of this study is to fundamentally quantify changes in strain associated with AI particle thermal treatments and examine energy propagation behavior of treated AI particles combined with CuO. These extensions enable an understanding of how thermal treatment and mechanical properties couple to influence the reactivity of AI particles with a solid oxidizer. Improving AI reactivity with a relatively simplistic heat treatment approach may greatly improve their functional application.

Experimental

Sample Preparation

Aluminum particles were procured from Alfa Aesar (Ward Hill, MA). Their average diameter was determined using an AccuSizer 780 optical particle size analyzer (Santa Barbara, CA). The Al particles were suspended in filtered water (specifically, 2 mg Al/80 ml water) and sonicated for 60 minutes in cycles of 10 seconds on, 10 seconds off in order to avoid heating. The sample was then pulled via a syringe pump for analysis. Particle length (diameter) is calculated and plotted as a count-based measurement. The plot for the untreated Al is shown in Fig. 25a and the size data is given in Fig. 25e. The oxide thickness used for purity calculations was estimated at 4 nm (i.e., >1% oxide concentration, see Fig. 24). The untreated Al particles have an average diameter of 5 microns.







Figure 25. Particle size distribution of (a) untreated AI and AI annealed to (b) 100 °C, (c) 200 °C, (d) 300 °C and (e) average particle diameter with standard deviation for each AI powder sample.

Because the oxide layer thickness is independent of Al particle size, the oxide thickness used for purity calculations was estimated at 4 nm (i.e., >1% oxide concentration(see Fig. 24)). The untreated Al particles had an average diameter of 5.9 microns.

Aluminum particles (200 mg) were loaded into ceramic trays and subjected to various annealing temperatures and quenched to room temperature. This process used a Neytech Qex vacuum oven (Torrance, CA). This is a programmable oven, such that the powder was annealed to 100, 200 or 300°C at a controlled rate of 10 KPM, held at the prescribed temperature for 15 minutes then removed from the oven and quenched via refrigeration to room temperature in air at standard pressure. Powder temperature was monitored using an InstruNet Direct to Sensor system (Charlestown, MA) and Type K thermocouples from Omega Engineering (Stamford, CT). Aluminum temperature was monitored for each thermal cycle and Fig. 26 shows the temperature response as a function of time.



Figure 26. Temperature plots for AI thermal cycles to (a) 100 °C, (b) 200 °C and (c) 300 °C.

For the natural convection conditions that exist in these experiments, materials cool according to a lump capacitance model. The quenching rate is an exponential function of time as shown in Eq. (16). The temperature evolution reduces exponentially and experimental results are approximated by Eq. (16). $T = T_a + (T_0 - T_a) \exp(-At) \text{ with } A = 0.0078 \text{ s}^{-1}$ (16) In Eq. (17) T_a is ambient temperature, T_0 is annealing temperature, t is time and A is determined by examining the exponential plots of the cooling curves and identifying the coefficient. An average quenching rate ranges from 0.13 to 0.38 Kelvin per second (KPS) depending on annealing temperature.

The Al particles were then mixed with 50 nm average particle diameter spherical CuO particles (Sigma Aldrich (St. Louis, MO)) to an equivalence ratio of 1.2 (i.e., slightly fuel rich). The mixing process is well documented(C. a. Crane, Pantoya, and Weeks 2014; Feng et al. 2013; K. Kappagantula, Crane, and Pantoya 2014) but will be summarized here. The dry powders were weighed and suspended in hexane and mixed using a Misonix Sonicator 3000 probe for 2 minutes. The solution is poured into a Pyrex dish and hexane evaporated in a fume hood for 24 hours. The dry powders were retrieved and sieved to break up large agglomerations.

The powder is carefully loaded into 3 mm inner diameter, 8 mm outer diameter, 10 cm long quartz tubes containing 850 mg of powder each. The theoretical maximum density (TMD) of the loose powder is determined by a weighted average of the bulk densities of AI (2.7 g/cm³), Al₂O₃ (3.95 g/cm³), and

CuO (6.31 g/cm³). This is calculated using Eq. (17) where *M* is the percent mass of reactant *i* and ρ is the density for each reactant.

$$\mathbf{TMD} = \frac{1}{\sum_{i} M_{i} \frac{1}{\rho_{i}}} \tag{17}$$

The calculated TMD is 5.46 g/cm³ and the bulk density is 1.04 g/cm³ such that tubes were loaded to 19% TMD.

Strain Measurements

X-Ray Diffraction experiments were performed at the Advanced Light Source on beamline 12.3.2 using a micron focused synchrotron x-ray beam. This is a unique facility that allows measurement of micron-scale samples. Aluminum powder samples were spread over glass slides and scanned under the x-ray beam (either polychromatic or monochromatic) while a diffraction pattern was collected at each step using a DECTRIS Pilatus 1 M detector. The measured relative small shifts in the reflection positions in the Laue pattern provides the deviatoric strain tensor of the material while the measurement of the energy of one reflection provides the dilatational component. Data were processed using XMAS software(Kunz et al. 2009). The beamline experimental setup and capabilities have been described elsewhere(Tamura 2013). Flame Speed Measurements

Figure 27 illustrates a typical powder filled tube arrangement for measuring flame speed and the apparatus is described in more detail elsewhere(Dikici, Pantoya, and Levitas 2010) but is summarized here.



Figure 27. Powder filled quartz tube and representative still frame images time stamped of powder filled quartz tube and flame propagation. Bulk density is 19% theoretical maximum density. Note nichrome wire extruding for left end of the tube.

Both ends of the tube were sealed with one side securing a length of nickel-chromium wire for ignition. Five experiments per annealing temperature were performed to establish repeatability. Each tube was placed inside a blast chamber for ignition and flame propagation experiments. The powders were ignited and flame propagation was observed through a viewing window in the chamber (as seen in Fig. 28). The reaction was recorded with a Phantom v7 (Vision Research, Wayne, NJ) high speed camera at a rate of 29,000 frames per second and 512 x 128 resolution. The camera was aligned perpendicular to the direction of flame propagation. Flame speed was determined by tracking the flame front through a referenced time and distance using the Vision Research Software. The resolution of the flame speed for this diagnostic is 0.1 m/s. The largest source of uncertainty in the measurement is due to repeatability and is shown for each data set in the results.



Figure 28. Ignition setup for tracking energy propagation.

Results

Further particle size analyses were performed for each Al sample after thermal treatment. Figures 25b-d show the plots for Al particle sizing and the average particle diameter and distribution is given in Fig. 25e. This data shows that the thermal treatments did not result in any sintering or agglomerations and that the average diameter is consistent (about 5.9 μ m).

Figure 29 shows results of dilatational strain (i.e. change in volume) distribution measurements of aluminum particles that were (a) untreated and annealing to (b) 100° C, (c) 200° C and (d) annealed to 300° C. Table 6 presents the count based averages for dilatational strain for each annealing temperature. The average strain for the samples annealed to 100 and 200° C fall within the resolution of the machine ($2x10^{-5}$), so their percent increase was negligible.

However, the aluminum particles annealed to 300°C and cooled to room temperature showed a significant dilatational strain increase from the baseline Al particles (i.e. 660%). It is noted that these measurements are of the aluminum core because the alumina shell is too thin (i.e., less than 5 nm) to resolve volume based strain measurements using this approach.



Figure 29. Dilatation strain for (a) untreated particles and particles annealed to (b) 100 °C, (c) 200 °C and (d) 300 °C.

Figure 30 shows flame speed as a function of annealing temperature. The samples annealed to 100°C experience a 2% increase in flame speed from the baseline; and, when annealed to 200°C flame speed increases by 3%. Once the

samples are annealed to 300°C, they experience a 23% increase in flame speed. This is a significant increase in reactivity for the composite annealed to 300°C but not for the other annealing temperatures. These results are also summarized in Table 6.



Figure 30. Flame speed as a function of annealing temperature. Average values for flame speed are reported above the bar and standard deviation in the measurements is also shown.

Table 6. Dilatational strain in Al particles, flame speeds when combined with CuO, and percent increase for all Al annealing temperatures. The average strain for the samples annealed to 100 and 200 °C fall within the resolution of the machine, so their percent increase was negligible.

Annealing	Average	%	Flame	%
Temperature	strain	increase	Speed	increase
(°C)			(m/s)	
untreated	1.00E-05		95.2	
100	1.47E-05		96.7	
200	1.53E-05		97.6	
300	3.25E-05	660	118	24

Discussion

Figure 29 shows that compared to the baseline, volumetric strain increases as annealing temperature increases. According to Fig. 29 and Table 6, average dilatational strain increases by 660% after thermal treatment to 300 °C but is unchanged in the samples annealed to 100 and 200 °C. Diffusional creep resistance occurs in samples when the annealing temperature is below a critical fraction of the melt temperature (i.e., T_m = 660 °C for Al)(Kitagawa, Jaske, and Morrow 1969). Kitagawa et al. investigated creep response in bulk Al samples subjected to a variety of temperatures (i.e., from $0.32T_m$ (~27 °C) to $0.55T_m$ (~240 °C)) for different number of thermal treatment cycles. They were interested to see the Average creep rate for each temperature over a variety of thermal cycles and found that the transition temperature for Al to exhibit creep acceleration is $0.4T_m$ (i.e., 100 °C) and the maximum acceleration of creep for Al occurs at about $0.52T_m$ (212 °C). Figure 29 shows that the samples annealed to 100 and 200°C show little response in terms of dilatational strain, consistent with annealing temperatures below the critical temperature for maximum acceleration of creep (i.e., 212 °C). However, once this critical temperature is passed (i.e., 300 °C), the Al particles responded with a 660% increase in dilatational strain. The effect of this trend is seen in the flame speed results. The Al samples annealed to 100 and

200 °C showed little change from the untreated case, but for the 300 °C case, flame speeds increased by 24%.

Ultimately, the annealing and quenching treatment effectively increased the overall strain. It is important to note that at higher annealing temperature or longer annealing times, particles could experience shell growth and/or oxide phase change. Amorphous alumina begins to transition to gamma-phase alumina around 440 °C, such that annealing beyond this temperature would affect the shell microstructure, which may also impact reactivity. Also, annealing to temperatures past 440 °C for an extended time (60+ minutes) would result in oxide layer growth as shown by Gesner et al.(Gesner, Pantoya, and Levitas 2012). They annealed 95 nm diameter Al particles to 480 °C for a variety of hold times and saw an increase in oxide shell thickness from 2.7 to 8.3 nm when held for 150 minutes but negligible shell growth for times applied in this study (and in this study annealing temperatures did not exceed 300 °C).

Thermal treatment is purposefully designed to relax residual stresses by annealing then reintroduce desirable stresses by quenching. While thermal treatments are relatively established for bulk metal processing, manipulation of mechanical properties for particles that consist of a core-shell structure are not well understood. In this study, results for thermal treatment of particles appear to be consistent with theory for bulk metals. The benefit of thermal treatment from Fig. 29 and 30 is that the annealed and quenched AI powders now have experience higher strain that also increases macroscopic energy propagation.

Conclusion

This study examines the effect of aluminum powder annealing and quenching treatments on the strain in of aluminum particles and corresponding reactivity. Micron-sized aluminum was annealed and quenched according to treatments designed to affect mechanical properties. Synchrotron x-ray diffraction (XRD) analysis of the particles reveals annealing increased the dilatational strain of the Al particles. Diffusional creep is the primary mechanism affecting strain and for powder annealed to 300 °C, a 660% increase in dilatational strain was observed. However, annealing to 100 or 200 °C showed no significant increase in dilatational strain. Treated Al powder was then mixed with CuO to assess reactivity. Flame speed measurements similarly show the 300°C annealed sample produces the highest flame speed. These results reveal that altering the mechanical properties of aluminum particles affects their reactivity, particularly when combined with a solid oxidizer.

OVERVIEW

Chapter two investigates the combustion performance of various Al particle sizes coated with PFPE (a liquid fluorinated oligomer). Flame speeds, differential scanning calorimetry (DSC), thermogravimetric analysis (TGA) and guadruple mass spectrometry (QMS) were performed for AI- PFPE blends with varying Al particle sizes (80, 100, 120 and 5500 nm in average diameter). The results show that the combustion performance of these blends is highly dependent on the Al₂O₃ concentration and exposed surface area. There is a balance to optimizing Al particle reactivity between activating Al particles with exothermic surface chemistry versus the unreacted alumina that contributes a thermal heat sink during energy generation. Overall, flame speed is inversely proportional to the apparent activation energy for the blend, with lower activation energy correlating with higher flame speeds. Also, combustion performance is optimized when the material concentrations are balanced such that they burn in a conductively driven manner.

Once fundamental combustion was understood for AI-PFPE blends, chapter three covered combustion characterization analyses for varying concentrations of PFPE in AI/MoO₃ and AI/CuO composites in order to examine the influence of fluorine surface chemistry on reactivity using high speed imaging, XPS, QMS, and DSC. Results show that the performance of the thermite-PFPE blends is highly dependent on the metal oxide. The PFPE blends with MoO₃ show an increase in reactivity while the blends with CuO show a decrease in reactivity when increasing PFPE concentration. We observed a decline in formation of AIF₃ in CBT samples but an increase in AIF₃ formation in the MBT samples. The potential surface chemistry between Al₂O₃ and F from PFPE is negated by competitive Al-O, but thermodynamically favorable formation from CuO with lower bond dissociation energy than F or the CF₂ radical. When using an oxidizer with a BDE similar to that of the C-C or C-F bond (i.e. MoO₃), PFPE can promote reactivity via catalytic behavior of the alumina shell to help decompose PFPE more efficiently and improve the low temperature surface chemistry and overall reactivity.

Chapter four examines the effect of aluminum powder annealing and quenching treatments on the strain in of aluminum particles and corresponding reactivity. Micron-sized aluminum was annealed and quenched according to treatments designed to affect mechanical properties. Synchrotron x-ray diffraction (XRD) analysis of the particles reveals annealing increased the dilatational strain of the Al particles. Diffusional creep is the primary mechanism affecting strain and for powder annealed to 300 °C, a 660% increase in

dilatational strain was observed. However, annealing to 100 or 200 °C showed no significant increase in dilatational strain. Treated Al powder was then mixed with CuO to assess reactivity. Flame speed measurements similarly show the 300°C annealed sample produces the highest flame speed. These results reveal that altering the mechanical properties of aluminum particles affects their reactivity, particularly when combined with a solid oxidizer.

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