SOLUTION ASSISTED LASER ABLATION SYNTHESIS OF DISCRETE ALUMINUM NANOPARTICLES (Preprint)

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Discrete aluminum nanoparticles with an average particle size of 21 nm have been prepared by laser ablation of a metallic aluminum target submerged in dry tetrahydrofuran in the presence of 0.001 M oleic acid as a stabilizing ligand. The particles display high solubility and minimal aggregation while the absence of oleic acid leads to highly aggregated particles and a broader particle size distribution. O/Al ratios obtained from EDS analysis of single particles and small particle clusters suggests the particles produced are primarily metallic aluminum with minimal oxide content.
Solution assisted laser ablation synthesis of discrete aluminum nanoparticles

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ABSTRACT

Discrete aluminum nanoparticles with an average particle size of 21 nm have been prepared by laser ablation of a metallic aluminum target submerged in dry tetrahydrofuran in the presence of 0.001 M oleic acid as a stabilizing ligand. The particles display high solubility and minimal aggregation while the absence of oleic acid leads to highly aggregated particles and a broader particle size distribution. O/Al ratios obtained from EDS analysis of single particles and small particle clusters suggests the particles produced are primarily metallic aluminum with minimal oxide content.

Keywords: Nanomaterials; metals and alloys; laser processing

1. Introduction

The high energy density and reactive nature of aluminum nanoparticles (nAl) has catalyzed interest into their use as fuel and/or additives in many propulsion and pyrotechnic formulations. The reaction kinetics for Al combustion are governed by surface diffusion, therefore transitioning from micron-scale powders to nanoparticles has lead to remarkable enhancements in reaction rates due to an exponential increase in the specific surface area of the reactant materials. To meet the increasing demand for nAl, many synthetic approaches have been explored including wet chemical methods, e.g.
catalyzed reduction of alane precursors [1,2,3] and hydrogenolysis of organometallic precursors [4], as well as physical techniques, *e.g.* electrical explosion of Al wire [5], plasma vaporization/inert gas condensation [6] and laser ablation [7]. The pyrophoric nature of nAl requires passivation with a thin (2-6 nm) amorphous oxide layer at the particle surface to serve as a barrier to further oxidation of the underlying metal. Though effective, this method consumes active Al and can result in a material with more than 40% unreactive oxide by weight, as the particle size drops below 30 nm [8]. To avoid loss of active Al due to oxide formation researchers have explored alternative passivation routes including the use of hydrocarbon [9] and perfluorinated ligands [3], polymers [9], and transition metal oxide coatings [8]. Though these approaches produce products which are relatively stable to environmental oxidation, they provide little control over particle size and/or yield a highly aggregated product. To further enhance the potential for use of these materials in propulsion and pyrotechnic applications it would be desirable to have a discrete product with minimal aggregation in addition to tunable size control.

Laser ablation in aqueous and organic solvents has regularly been used as a route for producing noble metal and metal oxide nanoparticles [10]. Particle size control is shown to be dependent upon operating conditions (wavelength, power, etc.) and thus a variety of particle size distributions can be achieved by manipulation of the experimental parameters. Additionally, stabilizing agents (*e.g.* polymers and/or surfactants) can be added to yield a stable colloid or to prevent/minimize aggregation in the reactant products [10,11]. Herein we report, to the best of our knowledge, the first successful example of solution assisted laser ablation to yield discrete, organically capped nAl from bulk metallic Al substrates in a dry organic solvent (tetrahydrofuran) containing a stabilizing ligand (oleic acid) to minimize particle aggregation.

2. Experimental

The experimental setup is illustrated in Fig. 1. A 250-mL four-neck tapered sonochemical flask (Ace Glass, Inc.) was charged with a one inch Al target and fitted with three rubber septa and a nylon
Oleic acid (OA)(tech. grade, >90%) and tetrahydrofuran (THF) (anhydrous) were obtained from Sigma-Aldrich and used as received. Aluminum targets (99.9995%, 1 in. diameter) were obtained from Kurt J. Lesker Company. The flask was evacuated and purged with dry nitrogen a minimum of three times to remove air and moisture. A 0.001 M solution of OA in anhydrous THF (ca. 25 mL) was added until the solvent level was ca. 1 cm above the surface of the aluminum target. The reaction vessel was then purged with nitrogen and sealed. A focusing lens (focal length = 300 mm) was positioned above the reaction vessel yielding a 1 mm spot size on the target. A 1064 nm Nd:YAG laser, operating at 20 Hz with a beam power of 0.88 J/pulse, was used to ablate the aluminum target for 5 minutes. The reaction mixture was transferred via canula to a sealed vial where it was stored under nitrogen prior to analysis. Transmission electron microscopy (TEM) samples were prepared by drop drying the solution onto a 200 mesh Cu grid coated with a holey carbon film (SPI, Inc.). TEM images were obtained on a FEI Titan microscope operating at 300 kV. Energy-dispersive X-ray spectroscopy (EDS) analysis was performed using an EDAX detector from 0-20 keV. Al$_2$O$_3$ nanoparticles (AMCN01, Accumet Materials Co.) were used as a standard for comparison of O/Al ratios. Particle size distribution data was obtained from diameter analysis of 471 particles from 10 separate TEM images and sample locations using ImageJ software.

3. Results and Discussion

Aluminum nanoparticles were prepared by solution assisted laser ablation of an Al target submerged in a 0.001 M OA solution in THF, and also in THF without any OA. As prepared these particles become suspended in solution and remain in solution for a period of several months, Fig. 2. TEM micrographs of the products are presented in Fig. 3. The OA stabilized nAl (OA-nAl) particles are noticeably discrete (non-aggregated) with a narrow size distribution displaying an average diameter of 21 nm as shown in the particle size distribution graph, Fig. 3(d). Some larger particles and fragments, 100 nm or more, are
also occasionally observed as shown in Fig. 3(b). Solubility of the product is in part due to the small particle size and their discrete nature but also the presence of an OA coating at the particle surface provides a hydrophobic shell which enhances miscibility with common organic solvents. As shown in Fig. 3(c), ablation in the absence of OA also leads to the formation of nAl, however, these particles are highly aggregated with only a small percentage of the particles remaining discrete.

EDS spectra, Fig. 4, were collected to analyze the O/Al ratios for the discrete OA-nAl. These ratios were compared to O/Al ratios obtain for commercial Al₂O₃ particles to qualitatively understand the chemical state of the OA-nAl. Spectra were acquired from both single particles and clusters of particles present on the TEM grids. Low O/Al ratios (Table 1), obtained from comparing the intensities of the aluminum signal (1.44 keV) to the oxygen (0.46 keV) signal, suggests that the particles mostly remain as metallic aluminum and are not fully oxidized during the synthesis or upon exposure to atmospheric conditions.

Previous reports on the preparation of nAl by laser ablation in a liquid phase [12] suggested the formation of nAl with an average particle size of 50 nm or less, however, no evidence was provided confirming the state of Al as either metallic or oxide. Moreover, the experimental design consisted of an ablation target formed from compressed Al powder with distilled water as the liquid phase. Due to the passivated oxide layer present on Al powders the ablation target would contain an elevated oxide content compared to a bulk metal target. Additionally, Al readily reacts with water to form aluminum oxide and oxide-hydroxides [13], therefore one might expect that the primary product would be aluminum oxide nanoparticles and not metallic nAl as suggested. To circumvent these complications in our studies we have chosen to perform all experiments under air-free conditions by operating with dry organic solvents under inert atmospheres. High purity Al sputtering targets were selected to minimize the presence of oxidized Al in the system. Additionally, bulk Al is perhaps the most stable starting material from which Al nanoparticles can be prepared, thereby eliminating the use of expensive and
sensitive powders and/or organometallic precursors. Previous literature reports have utilized the
coordination of various carboxylic acid ligands to the metallic Al surface to prevent oxide passivation of
the particles [3,14]. With similar intentions, OA was added to our reaction mixture in an attempt to
inhibit oxidation of the produced metallic nAl through formation of an organic layer at the particle
surface immediately upon particle formation. The presence of OA additionally aides in minimizing
aggregation and promotes solubility of the nanoparticle product in the reactant solution. EDS spectra of
the OA-nAl demonstrate relatively low oxygen content, which can be partially attributed to oxygen
contained within the OA ligand. XPS analysis does confirm the presence of oxidized Al suggesting that an
oxide layer is still present or that some of the smaller particles are partially oxidized. It is unclear
whether this oxidation occurs during the reaction or upon sample preparation and exposure to the
atmosphere but in either case OA was not completely successful at inhibiting oxidation although it does
prevent aggregation. Similar results have been reported for the sonochemical synthesis of Al
nanoparticles in the presence of an OA/dodecane mixture [14].

4. Summary

Discrete metallic nAl have been successfully prepared by laser ablation of an Al target in the presence of
a dry organic solvent, THF. Aggregation of the particles was minimized through the addition of OA,
whereas particles produced in the absence of OA readily from aggregates with only a small amount of
particles remaining discrete. OA also appears to impart some control over the average particle size of
the product leading to a narrow, unimodal size distribution displaying an average particle size of 21 nm.
The discrete nature and small particle size allow for the preparation of stable particle suspensions in THF
with the particles remaining suspended for months. This work provides a novel and unique route
towards the synthesis of stable, discrete metallic nAl from stable and inexpensive Al precursors.

Acknowledgment
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Figures

![Schematic of the experimental setup](image)

**Fig. 1.** Schematic of the experimental setup.
Fig. 2. Digital images of the reaction taken a) before and b) after ablation (5 minutes).
Fig. 3. TEM micrographs of (a,b) discrete OA-nAl, c) nAl produced in THF without OA and, d) particle size distribution for OA-nAl.
**Fig. 4.** EDS spectra of a) discrete OA-nAl nanoparticles and b) Al$_2$O$_3$ nanopowder (control).

**Table 1** O/Al ratios obtained from EDS analysis of OA-nAl and Al$_2$O$_3$ nanoparticles

<table>
<thead>
<tr>
<th>Sample</th>
<th>O/Al</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single particle</td>
<td>0.094</td>
</tr>
<tr>
<td>Clustered particles</td>
<td>0.159</td>
</tr>
<tr>
<td>Al$_2$O$_3$ (control)</td>
<td>0.603</td>
</tr>
</tbody>
</table>
References