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Effects of Oxidation on the Plasmonic Properties of Aluminum Nanoclusters†

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The scouting of alternative plasmonic materials able to enhance and extend the optical properties of noble metal nanostructures is on the rise. Aluminum is endowed with a set of interesting properties that turn it into an attractive plasmonic material. Here we present the optical and electronic features of different aluminum nanostructures stemming from a multilevel computational study. Molecular Dynamics (MD) simulations using a reactive force field (ReaxFF), carefully validated with Density Functional Theory (DFT), were employed to mimic the oxidation of icosahedral aluminum nanoclusters. Resulting structures with different oxidation degrees were then studied through the Time-Dependent Density Functional Tight Binding (TD-DFTB) method. A similar approach was used on aluminum nanoclusters with a disordered structure to study how the loss of crystallinity affects the optical properties. To the best of our knowledge, this is the first report that addresses this issue from fully atomistic time-dependent approach by means of two different and powerful simulations tools able to describe quantum and physicochemical properties associated with nanostructured particles.

1 Introduction

The study of plasmonic properties of very small metallic particles have been mainly focused on gold and silver nanostructures. Colloidal solutions of silver and gold nanostructures often exhibit an intense optical absorption band located on the visible region of the electromagnetic spectrum owing to coherent oscillations of conduction electrons in response to external electric fields. These optical features along with their noble character render silver and gold nanostructures the ideal plasmonic metals. Nowadays, there is an extensive research of diverse materials that exhibit optical responses similar to noble metal nanostructures but with a plasmonic excitation located beyond the visible region. In this context, aluminum appears as one of the most promising plasmonic materials. Not only it exhibits attractive plasmonic properties, but it is also biocompatible, abundant and a low-cost material.

The plasmonic properties associated with very small aluminum nanostructures are characterized by having a single dipole surface plasmon resonance (SPR) band located below 400 nm within the ultraviolet region. Thereby, the plasmonic response of aluminum nanoclusters would fill up a region of the spectrum where the optical properties of silver and gold nanostructures are intrinsically inactive. This distinctive optical feature would allow electronic coupling between plasmonic excitations and HOMO-LUMO transitions (220-280 nm) for a large number of bio- and organic molecules.

From the synthetic point of view, the production of colloidal aluminum nanoclusters have been scarcely explored and only a few methods based on wet chemical have been proposed. Despite its promising plasmonic properties, aluminum is well known for spontaneously form a thin oxide layer in presence of O2. The oxide layer prevents the metal for further oxidation, a phenomenon known as passivation. Recently, a synthetic pathway for formation of bare aluminum nanoclusters in organic media has been reported. In regular conditions, however, the formation of an oxide layer is inevitable. For nanostructured aluminum this oxidation process may modify its inherent plasmonic properties.

Theoretical studies of optical properties associated with different metallic nanostructures have been carried out through the use of several models and computational tools. Within the framework of methods based on classical electrodynamics, the discrete dipole approximation (DDA) has been one of the most widely used to explore and predict different optical phenomenon associated with particles with diverse compositions and morphologies.

† Electronic Supplementary Information (ESI) available: Validation of the Al-O ReaxFF from comparison with results obtained from DFT for small Al2O3 clusters. See DOI: 10.1039/b000000x/
However, when the particle size is reduced (diameters of less than 10 nm) several size-related surface and quantum effects start to govern optical response and the DDA starts to lose validity. In this size domain, the optical response of metallic nanostructures can be obtained from methods based on time-dependent density functional theory (TD-DFT) approach. This approach includes all factors that determine the optical properties in a quantum regime (depending on the nature of the exchange-correlational functional used). Despite nowadays being the “weapon of choice” to study the absorption spectra of molecular systems, the method has a strong limitation with respect to the size of the approachable systems. This limitation is essentially related with computational cost. In recent years, many systems including silver\textsuperscript{18–26}, gold\textsuperscript{21,22,27–34}, aluminum\textsuperscript{35,36} and bimetallic\textsuperscript{32,37–40} nanostructures have been explored by using TD-DFT.

The optical properties of different aluminum nanostructures have been recently investigated using mainly the discrete dipole approximation (DDA) method.\textsuperscript{9–11} Likewise, the effect of aluminum oxide over plasmonic properties have been studied using the same classical electrodynamics methods\textsuperscript{10}. However, this type of approximation does not take into account the complex size-related-quantum effects that govern the optical properties for particles sizes below of 5 nm, nor the influence of the oxide layer on the electronic structure of the aluminum core. Despite the abundant research on the subject, the effect of oxygen on the optical properties of these nanostructures has not been undertaken from a fully atomistic approach. In order to do so, we have performed a multi-level systematic study combining Molecular Dynamics (MD), Density Functional Theory (DFT), and Quantum Dynamics. The optical absorption properties of bare icosahedral aluminum from 1.1 nm to 3.3 nm in diameter were explored and compared with those of the corresponding oxidized nanostructures.

## 2 Computational Method

### 2.1 Molecular Dynamics Simulations

Energy minimizations, molecular dynamics simulations, and simulated annealing were carried out using the Large-scale Atomic-Molecular Massively Parallel Simulator (LAMMPS)\textsuperscript{41} with its ReaxFF module,\textsuperscript{42} which describes all the interactions among Al and O atoms.\textsuperscript{43} Cubic simulation cells were used for nanoclusters at 1, 12, and 160 atm of O\textsubscript{2} pressure for which the corresponding side lengths were 298, 138, and 65 Å. As the oxidation reaction proceeds, the simulation box shrinks in order to keep the pressure constant. We carefully checked that the gap between a nanocluster and its periodic images was greater than 20 Å during the whole simulation in order to avoid self-interactions. Initial velocities were randomly assigned following a Maxwell-Boltzmann distribution. Periodic boundary conditions were used in three dimensions.

The nanoclusters were subject to three different O\textsubscript{2} pressures: 1, 12, and 160 atm in an NPT regime (constant number of atoms, pressure, and temperature) with a time-step of 0.2 fs using the Nose-Hoover algorithm. This entails that as the O\textsubscript{2} reacts with the metallic nanocluster the volume of the simulation cell decreases so as to keep the pressure constant. This scenario is more realistic than the NVT regime (constant number of atoms, volume, and temperature), in which the oxygen pressure decreases as the reaction proceeds.

It was found that reaction speed is proportional to oxygen pressure. Indeed, at the lowest pressure value (1 atm) the reaction is so slow that it becomes infeasible for a systematic study within our simulation approach. At intermediate oxygen pressures (12 atm) the nanocluster suffers fragmentation, which also make the structures useless for further studies. This phenomenon of oxide cluster formation with a variety of aluminum-oxygen ratios have been found before in both simulations,\textsuperscript{44,45} and experiments.\textsuperscript{46,47} Fragmentation was also found at 1 atm of O\textsubscript{2} pressure. Details concerning geometry configurations and temperature of the corresponding nanoclusters are given in the SI. Finally, at 160 atm, the oxidation is fast and without any sign of fragmentation, we therefore chose this set up for the following calculations as a means of obtaining passivated aluminum cluster structures. This value of pressure corresponds to the lowest value studied in ref.\textsuperscript{43}

Given that the most essential part of a MD simulation is the force field used, we dedicate a few lines to explain its validation. The Al-O ReaxFF has been proven to qualitatively predict the oxidation kinetics of aluminum slabs and nanoclusters as a function of temperature.\textsuperscript{43} In order to take the validation one step further, we performed DFT relaxations of small Al\textsubscript{2}O\textsubscript{4} clusters, and compared the resulting structures and energies with that obtained by the ReaxFF. This study showed that ReaxFF favors the coordination of oxygen with four Al neighbors (inside an aluminum tetrahedron), while for DFT three Al neighbors is energetically more stable (inside an aluminum triangle). Also, the force field tends to overestimate the formation energy (see eq. 2). However, the energy discrepancy diminishes as the cluster becomes larger, where quantum effects are less accentuated. Despite these deviations, ReaxFF correctly estimates the relative energy among structures (see the supporting information for details).

For the simulated annealing calculations the oxidized aluminum nanocluster were first relaxed, then the temperature was increased from 0 K to 800 K, and then decreased to 0 K in an NVT ensemble using a Nose-Hoover thermostat during 200 ps. In order to analyze the degree of oxidation of such structures, the same criteria as in ref.\textsuperscript{43} was used. Aluminum atoms are classified according to the percentage of metallic neighbors up to a radius of 3.2 Å, they are considered as metal (more than 90%), suboxide (between 60% and 90%), oxide (between 40% and 60%), peroxide (less than 40%). Then, each aluminum atom is counted between two radial distances \( r \) and \( r + dr \) from the central atom (the atom closer to the nanocluster center of mass). To cancel out the increase of shell volume with \( r \), this count is divided by \( N(r) \), which is the shell volume \( (V_s) \) times the density of the Al fcc bulk \( (\rho) \).

\[
N(r) = \frac{n}{V_s \rho} \tag{1}
\]

For comparison of stabilities the formation energy was calculated as

\[
E_f = \frac{E(n)_{\text{AlO}} - E_{\text{Al}} - nE_{O_2}}{n} \tag{2}
\]
where $E(n)_{AIO}$ is the energy of an oxide cluster with $n$ number of oxygen atoms, $E_A$ the energy of a bare aluminum cluster, and $E_O$, the energy of an oxygen molecule.

2.2 Density Functional Theory

Density Functional Theory (DFT) calculations were performed using the Quantum Espresso/PWSCF code.\(^{48}\) Vanderbilt ultrasoft pseudopotentials\(^{49}\) were used together with the Perdew-Burke-Ernzerhof (PBE)\(^{50}\) approximation to the exchange-correlation functional. A 30 Ry kinetic energy cutoff and a 300 Ry charge density cutoff where used. Only the gamma point was considered in the electronic structure calculations owing to the finite nature of the system. Geometry relaxations were converged when forces were less than 0.02 eV/Å. Unit cells were big enough so that clusters were separated by at least 12 Å from their periodic images. DFT calculations where used to validate the Reax Force Field. We performed DFT relaxations of small Al$_2$O$_3$ clusters, and compared the resulting structures and energies with that obtained by the ReaxPF. The results of these calculations are shown in the Supporting Information.

2.3 Time Dependent Density Functional Tight Binding

All spectra and plasmon lifetimes where calculated in the framework of the self-consistent-charge density-functional-tight-binding (SCC-DFTB) method.\(^{51-53}\) This method is based on a controlled approximation of density functional theory (DFT) and has been successfully employed to describe the electronic structure and quantum properties of large organic and bio-inorganic systems.\(^{53,54}\) The DFTB Hamiltonian is based on a second order expansion of the the full Kohn-Sham energy in the neighborhood of a reference density consisting of the superposition of atomic densities up to second order. The exact second order expansion is then approximated as the sum of a band energy term, a self-consistent energy term consisting of the damping electrostatic interaction between Mulliken charges and a local Hubbard like term plus an empirical repulsive term. All three body center integrals in the energy functional expression are neglected and compensated by the empirical repulsion. All two body Hamiltonian and Overlap matrix elements are explicitly calculated using a pseudopotential to represent core electrons and a minimal valence basis consisting of Slater type orbitals. For all time-dependent calculations, since the nuclei are frozen, only the exactly calculated terms appear and the sole approximations are the self-consistent Hamiltonian component as well as the neglect of three center integrals.\(^{51}\)

The electronic structure of the ground state (GS) in all studied systems, was calculated using the DFTB+ package,\(^{55}\) which is an implementation of the SCC-DFTB method. The DFTB+ code is used to compute the GS Hamiltonian (H$_{GS}$) and overlap ma-trix (S) and thus to obtain the initial GS reduced single-electron density matrix ($\rho$). The matsci-0.3\(^{56,57}\) DFTB parameter set was employed in order to obtain the electronic structure for both bare aluminum and aluminum oxide nanoclusters. This parameter set includes $d$ orbitals for aluminum atoms, extending the minimal basis set normally used in DFTB.

The methodology applied to describe electronic dynamics of the whole system is based on the real-time propagation of the reduced single-electron density matrix ($\rho$) under the influence of an time-varying external potential. This approximation is an extension of SCC-DFTB model to the time domain and allow us to obtain valuable information about electronic dynamics triggered by an external perturbation. The simulations all proceed through the numerical integration of the single electron density matrix equation of motion (EOM) within the SCC-DFTB model. This equation of motion is non-linear, since the Hamiltonian depends on the density matrix via the self-consistent terms and includes the same physics, albeit approximately, as the full fledged TD-DFT framework using a time local exchange correlation functional with the advantage that it can be applied to much larger systems using relatively modest computational resources. Our implementation uses General Purpose Graphical Computing Processors (GP-GPUs) in order to accelerate the computational integration of the EOM. The method has been described in detail in\(^{58,59}\).

The results shown correspond to two types different types of external time-varying potential. For the calculation of absorption spectra, the external potential corresponds to a Dirac Delta potential applied $t = 0^-$. For this calculations we use a scheme equivalent to that originally proposed by Yabana and Bertsh.\(^{60}\) After application of the perturbation the evolution of the density matrix is used to obtain the time dependent dipole moment. The Dirac Delta potential intensity ensures that the system evolves within the linear response regime. The conditions for this are described in Yabana and Bertsh’s work.\(^{60}\) Absorption spectra are obtained after a Fourier transform of the dipole signal and deconvolution, which in the case of a Dirac Delta perturbation amounts to a mere multiplication. In order to adequately deal with the time over which the EOM is evolved, the time dependent dipole moment signal is exponentially damped using a time constant of 10 fs. This amounts to adding a homogeneous broadening to all spectral lines of 0.033 eV.

Plasmon lifetimes can in principle be obtained directly from the spectra, as has been shown in the literature\(^{61,62}\), by fitting the plasmon peak to a Lorentzian lineshape. We have found, however, that from a time dependent perspective it is more convenient and reproducible to calculate the lifetime directly from the dipolar signal damping in response to a monochromatic continuous perturbation. The intensity of this perturbing signal being equivalent to the one described in the previous paragraph to ensure that the system evolves within the linear response regime. We have used this method in order to describe the Chemical Interface Damping caused by molecular adsorption on silver clusters.\(^{63}\) Working within the linear response framework, our method describes the plasmon lifetime arising from pure dephasing of excitations close in energy to the plasmon peak. This method is useful for large particles in which a saturation of the dipole signal is observed, i.e. a quasi stationary state is reached.

3 Results and discussion

This section is divided in three parts: first, the optical properties of different bare icosahedral aluminum nanoclusters are analyzed; then, the molecular dynamics procedure used to build
oxidized structures from the bare aluminum nanostructures is shown; and finally, the influence of oxygen on the optical absorption properties are revealed.

3.1 Bare icosahedral Aluminum Nanoclusters

Gold, silver, and aluminum present a face centered cubic (fcc) crystalline lattice, their lattice constant being 4.08 Å, 4.09 Å, and 4.05 Å, respectively. Several structures were studied for all the nanocluster sizes considered in this study, in each case we found that the icosahedral structure is the most stable. Details regarding geometries and energetics are given in the SI.. In this report, different bare icosahedral aluminum nanoclusters, ranging from 1.1 nm to 3.3 nm, were considered. A schematic representation of the aluminum nanostructures studied are shown in Figure 1.

Aluminum belongs to the boron group, its valence electrons occupy only s- and p- orbital shells. However, in our calculation, the d-electron orbital shell was also included in order to achieve a suitable description of aluminum optical and structural properties. Figure 2a shows the optical absorption spectra corresponding to the aluminum nanostructures shown in Figure 1. All the spectra are characterized by the presence of a single dipole surface plasmon resonance (SPR) band located near on ~ 5 eV (~ 248 nm) in the ultraviolet region in agreement with previous reports9–11,13. The fact that a single dipolar SPR band is observed is to be expected given the size of the particles, precluding multipolar bands to show any appreciable cross section, and in accordance to their symmetric shape that does not allow a geometric splitting of the band as would be the case for rod like particles, for example. An increase in intensity of dipole SPR band is simultaneously detected with a small redshift of plasmonic excitation energy when the particle size increases in agreement with other experimental12 and theoretical9–11 reports.

The plasmonic properties of bare aluminum nanoclusters resemble that of silver nanoclusters9. The optical absorption spectra of both metals are characterized by a single dipole SPR band without a detectable contribution of interband electronic transitions to the optical absorption spectrum. However, in the aluminum case, the redshift is less pronounced and less sensitive to particle size than in the case of silver. The density of states (DOS) and projected density of states (pDOS) over electron orbital shells of a bare icosahedral nanocluster (Al961) are shown in Figure 2b. It is important to stress that the sp-band spreads through the whole electronic energy window; from occupied to unoccupied states. On the other hand, the d-band is mainly composed by unoccupied states (above the Fermi energy).

In order to explore the underlying electron dynamics triggered by the light absorption process, a bare icosahedral Al961 nanocluster is illuminated with a frequency in tune with the excitation energy of the SPR band as is outlined in the Figure 2d. The excited electron dynamics at the frequency of plasmonic excitation (Figure 2a, blue solid line, 5.03 eV or 246 nm) is shown in the Figure 2c. The electron dynamics is characterized by the depopulation of molecular states below Fermi energy, states associated to the sp-band (green and blue lines), and the population of unoccupied molecular states mainly associated with the d-band (red line). This dynamical picture captures all the electronic transitions that are related with plasmonic excitation. In this sense, the optical response of aluminum resembles that of silver since both show collective intraband transitions. The time dependent populations for a range of energies are shown in Figure 2d. These are obtained from projecting the instantaneous density matrix onto the ground state orbitals. The populations show oscillations in tune with the plasmon frequency for a range of energies, in agreement with results by Townsend et al64.

The time dependent dipole moment (µ(t)) obtained from an illumination simulation is a direct emergent of the electron dynamics and can be used to determine both the homogeneous line width (Γ) and the related lifetime (τ) associated with the plasmonic excitation. This approach have been recently employed by us to determine the homogeneous line width associated with dipole SPR band for different capped and naked silver nanoclusters.65 Here, the same model that relate the time-dependent dipole moment with homogeneous line width has been adopted obtaining the following equation

\[ \Lambda(\tau, \Gamma) = k_1 \{ \exp\left(\frac{-\Gamma t}{2}\right) - 1\} + k_2 \{ \exp\left(\frac{-\Gamma t}{2}\right) + 1\} \]  

(3)

where \(k_1\) and \(k_2\) are constant independent of \(\Gamma\) and are related to the absorption coefficient and electric field strength. The obtained dipole moment for aluminum and silver nanoclusters with 561 atoms and its respective fitting are shown in Figure 3. The homogeneous line width and lifetime associated with plasmonic excitation for both metals are summarized in the Table 1. As it can be seen, marked differences exist between both dipole moment profiles. The homogeneous line width associated with the Al961 nanocluster (\(\Gamma_{Al961}\)) is larger than the corresponding to Ag561 nanocluster (\(\Gamma_{Ag561}\)). This difference means that the natural mechanisms of plasmon dephasing are increased in the aluminum case. Likewise, the spectroscopic feature associated with dipole SPR band can be characterized by other important parameter, the quality factor(Q). In this case, \(Q_{Ag}\) is larger than \(Q_{Al}\) in agreement with other theoretical reports.9
3.2 Aluminum Oxide Nanoclusters

Molecular dynamics simulations were performed for several aluminum nanoclusters at 300 K and 160 atm of oxygen pressure in an NPT ensemble. The evolution of the reaction was followed by plotting both the percentage of dissociated O$_2$ and the percentage of pure Al atoms during the simulation, as shown in Fig. 4a. The equilibrium of the oxidation process at this pressure is reached in the order of 40 ps. At this point there is still $\sim$70% of O$_2$ in the gas phase and $\sim$8% of metallic Al in the cluster. These are indications of a passivation effect in which the oxide formed prevents the oxidation from continuing. As explained above, the high pressure at which the aluminum nanoclusters is under was chosen in order to accelerate the reaction. However, this may entail that the oxidized cluster is trapped in a configuration far from that of minimum energy. It is reasonable to expect that at lower oxygen pressure (and therefore lower oxidation rate) the NP would have longer time to explore the potential energy surface and find lower energy configurations. In order to circumvent this problem, simulated annealing was performed on three stages of oxidation: A, B, and C, shown in Fig. 4a. This procedure allows the struc-
while on stage C they become significant and appear at distances from the central atom at which the metallic species disappear.

3.3 Electronic vs geometric effect

The optical absorption spectra of structures A, B, and C are shown in Fig. 5c. An increase in thickness of the oxide layer produces an optical response that is characterized by red-shift, a broadening and decrease in the intensity of dipole SPR band associated with aluminum nanostructures. This optical behavior is in good agreement with other theoretical and experimental reports. Halsas et al report how an oxide layer lowers the energy and intensity of the dipolar plasmon resonance in Al nanodisks. The same effect is reported from theoretical calculations based on the discrete dipole approximation by Zeng et al. Reports in the literature are for very large structures and therefore well described by classical models. Our results, despite showing the same qualitative features, show that these carry on down to very small sizes and resemble the main spectroscopic changes that characterize chemical interface damping. Our calculations extend the results present in the literature to a regime where the atomic structure comes into play and allow a deeper insight into the chemical effects on the plasmon lifetime and their relation to disorder.

The most intuitive interpretation of these results is that oxygen is directly affecting the plasmonic properties of aluminum. However, one could argue that it is the breakage of the highly symmetric structure what weakens the plasmonic signal, in which case the oxide would be affecting the plasmon indirectly. In order to determine which effect is dominant, a totally random structure of bare Al561 was created. A comparison of the electronic properties of the latter with that of a perfect icosahedral Al561 should manifest the geometric effect. Icosahedral and random Al561 were thermalized at 300 K during 200 ps. The latter was created by taking away the oxygen atoms of structure C and then thermalizing. Fig 6a shows the resulting structures. It can be seen that the icosahedral structure retains its geometry during the simulation. The radial distribution function g(r) of the metallic nanoclusters is given in Fig 6b. As expected, peaks of the random structure are wider and lower with respect to those of the icosahedral structure, indicating a larger deviation from the atomic crystal positions. Moreover, the peak of second neighbors found in the icosahedral structure is missing in the random structure.

As shown by the optical absorption spectrum of icosahedral and random Al561 in Fig. 6c, the loss of symmetry only produces mild modifications over the optical response. This result determines that the weakening of the plasmonic signal on AlO nanoclusters can only be attributed to the electronic effect induced by oxidation. A deeper insight over the effect of oxide formation can be obtained analyzing the projected density of state and electron dynamics, Fig 7. As it can be seen in the a), c) and d) panels of Fig. 7, the increase in thickness of the aluminum oxide layer gradually produces a separation of the continuous and featureless metallic density of states of the bare cluster (shown in Fig. 2b) into what constitutes the seeds of the conduction (lower energy) and valence (higher energy) bands of the pure oxide. A gap of reduced density of states around 5 eV below the Fermi level can be
clearly seen to deepen upon increasing oxidation. The lower energy (valence) band is mainly composed by oxygen orbitals while the lower energy (conduction) band is mainly composed by aluminum orbitals. There is a significant change at the density of states at the Fermi level. Whereas in the bare metallic cluster the DOS is monotonically increasing at the Fermi level, a peak is formed upon oxidation, that sharpens with increasing oxidation. Both the formation of a pseudo-gap below the Fermi level and the changes in the DOS at the Fermi level point to a diminishing of the metallic character of the particle. This weakening of metallic character induces more attenuated electron dynamics as can be seen in the panel e), f) and g) inside of Fig.7.

4 Conclusion

We have presented results of molecular dynamics simulations with a reactive force field combined with time-dependent density functional tight binding (TD-DFTB) simulations to describe the oxidation process of bare icosahedral aluminum nanoclusters and to explore the modifications that this oxidation causes on their optical absorption properties. It was found that oxygen induces a redshift and a broadening of the dipole surface plasmon resonance (SPR) band, which is in good agreement with both theoretical and experimental reports. These effects are only attributed to oxidation and not to the loss of the crystalline structure caused by the oxidative process. To the best of our knowledge, this is the first report that addresses the issue of the modification of optical properties of sizable metallic clusters due to oxidation through the combination of a series of atomistic methods that go from classical to quantum dynamics taking fully into account the atomic structure of the system.
Fig. 7 a), b) and c): Total density of states (black dot) and projected density of state over aluminum (gray solid line) and oxygen (red solid line) electron orbital shells for the corresponding oxidized structures A, B, and C; d), e) and f): Electron orbital population dynamics resulting from continuous light irradiation at dipole SPR band for the corresponding oxidized structures A, B, and C.

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