Computational prediction of a $+4$ oxidation state in Au via compressed AuO$_2$ compound

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Abstract
Much effort has been devoted to the investigation of the physical and chemical properties of the Au–O system over a range of pressures, owing to the considerable importance of these materials in fundamental and practical applications. To date, however, only Au$^{1+}$, Au$^{2+}$, Au$^{3+}$, and Au$^{5+}$ oxidation states have been identified in the Au–O system, but tetravalent Au$^{4+}$ has not been found. Here, we report the results of structure prediction for the Au–O system at high pressure via the effective structure prediction methodology within a first-principles electronic structure framework. We have uncovered an intriguing structure with AuO$_2$ composition and tetravalent Au, stable at high pressures. This phase shows an electronic transition from a metal to a semiconducting phase as a function of pressure. The present results provide fundamental understanding of the structural and physicochemical properties of compressed Au–O compounds.

Keywords: structure prediction, AuO$_2$, oxidation states, tetravalent

(Some figures may appear in colour only in the online journal)

Introduction
Gold (Au), being a well-known noble metal, is easily alloyed with other metals, but rarely reacts with other non-metal elements to form compounds [1, 2]. Therefore, much effort has been devoted to exploring the possible existence of unusual valence states for gold. For example, previous theoretical studies have revealed that Au may exhibit valence states ranging from $-1$ to $+6$ by combination with elemental lithium or fluorine at high pressures [3, 4]. However, it is well known that lithium and fluorine are extremely difficult to be prepared in high-pressure experiments. In the search for alternative oxidation states and compounds, the focus has shifted to other potential ligands, including oxygen, which has a high electronegativity value and is relatively easy to handle within the high-pressure experiments.

The most stable structure of gold oxide is Au$_2$O$_3$ ($Z = 8$, space symmetry group $Fdd2$) which is found to be stable at ambient conditions [5–7]. Orthorhombic Au$_2$O$_3$ contains 40 atoms per cell in the form of a network of AuO$_4$ units, including two crystallographical distinct types of O atoms and one type of Au atom. This Au$_2$O$_3$ compound is a semiconductor with a band gap of about 0.85 eV [5], but it is found to be unstable at high temperature [8]. Au$_2$O has also been experimentally identified as a breakdown product of Au$_2$O$_3$ when it becomes unstable under thermal treatment [8]. High pressure is known...
to be an effective tool to discover new phases with interesting physicochemical properties [9]. A recent high-pressure study [10] reported some new structures of AuO, which contain mixed-valent Au$^{3+}$ and Au$^{4+}$, as determined from the Bader and projected density of electronic states (PDOS) analysis within PBEsol functional [10]. A similar set of valence states are also reported in a previous study of Ag$^{1+}$/Ag$^{3+}$O compounds [11]. In a further piece of work [12], a new compound of AuO$_2$ was identified (space group $C2/m$, $Z = 4$) as stable at 50 GPa, in which Au is mixed-valence Au$^{3+}$/Au$^{5+}$ as computed from the Perdew–Burke–Ernzerhof (PBE) functional. Thus far, the $+4$ oxidation state of Au has not been found in Au oxides, but only in AuF$_4$ compounds as predicted under high pressure [3].

Here, we report the results of structure searches for the possible candidates of oxygen-rich compounds in the Au–O system, aiming to identify unusual Au$^{4+}$. Our results based on the global minimization of free energy surfaces merging $ab$ initio total-energy calculations through CALYPSO [13, 14] at high pressure provide useful guidance for further experimental synthesis. We found that a $Pb$-3 phase is much more stable than the previously predicted phase [12], indicating Au may have a $+4$ valence state within the Au–O compounds under pressure. At elevated pressure, it is found that metallic $Pb$-3 AuO$_2$ transforms to a semiconducting $C2/m$ phase. According to enthalpy calculations, we propose that the AuO$_2$ compound with Au$^{4+}$ is expected to form in experiments via the reaction of Au$_2$O$_3$ + O$_2$ or Au + O$_2$ at high pressure and high temperature.

### Computational details

We carried out systematic structure searches of the Au–O system at 0, 25, 50, and 100 GPa via the swarm-intelligence based CALYPSO code, which has been widely and reliably employed in the exploration of a large number of high-pressure systems [15–19]. In structure predictions, each generation contained at least 30 structures, and the first generation was produced randomly with symmetry constraints. The 60% lowest-enthalpy structures from each generation were used to produce the structures in the next generation, using the local particle swarm optimization (PSO) technique, and the remaining 40% of the structures were randomly generated within symmetry constraints to enhance the structural diversity. The CALYPSO method performs global structural minimization of free-energy surfaces combining density functional theory (DFT) calculations with the Vienna $ab$ initio simulation package (VASP) [20] code, unbiased by any known structure information. The PBE [21] and PBEsol [22, 23] were adopted for the exchange-correlation functionals. The electron-ion interaction was described using the all-electron projector augmented-wave (PAW) method [24] and $5d^{10}6s^1$ and $2e^22p^4$ were treated as valence electrons for Au and O, respectively. A plane-wave kinetic energy cutoff of 520 eV and dense $k$-point sampling by Monkhorst–Pack scheme with a grid of 0.025 Å$^{-1}$ were chosen to ensure that enthalpy calculations converged well. We employed the PHONOPY code [24] to calculate the phonon dispersion curves. Bader’s quantum theory [25] of atoms in molecules was used for the analysis of charge populations.

### Results and discussion

We systematically investigated the chemical stabilities of the Au–O system with 1–4 formula units per unit cell via the formation enthalpies at 0 K, relative to the products of their decomposition into constituent elements (Au and O), as summarized in figure 1. The PBEsol functional is known as a reparameterization of the PBE functional, which is suitable for solids and generally gives better description of experimental lattice constants, elastic properties, and cohesive energies [10, 22, 23]. We thus calculated the lattice constants of Au$_2$O$_3$ within PBEsol and PBE functionals at 0 GPa as shown in table 1, where our results are compared with the experimental observations [7]. The differences between the experimental data and PBEsol results for the lattice constants of Au$_2$O$_3$ are less than 0.5%, as shown in table 1. It is clearly seen that the PBEsol results are in better agreement with experimental data than the PBE results. The rest of our results are thus calculated within PBEsol functional in this work. Our calculations indicate that $Fd3d$ phase of Au$_2$O$_3$ is the ground state, in agreement with experimental observations [7], which validates the reliability of our computational scheme. The results indicate that Au$_2$O$_3$ gradually becomes unstable as pressure increases, until AuO$_2$ emerges as the most stable compound, as seen by the fact that it lies on the convex hull shown in figure 1.

![Figure 1. Calculated enthalpy of formation per atom of the Au–O system with respect to decomposition into elemental Au and O. The dashed lines connect the data points, and the solid lines represent the convex hull, where stable phases of O and Au are predicted at different pressures.](image-url)
In our structure prediction of the unconventional Au–O compounds, we found two stable phases for AuO2 with space group $\text{Pa}-3$ and $\text{C2/m}$, respectively (table 2). The $\text{C2/m}$ phase has been reported previously [12]. We find that the highly symmetric $\text{Pa}-3$ phase of AuO2 is stable between 42 and 50 GPa (figure 2). The $\text{Pa}-3$ phase takes a lattice constant of $a = 4.822$ Å (figure 3(a)), which is similar to the pyrite (FeS2) phase [26]. In this structure, Au and O lie on Wyckoff positions $\text{O1}(0.000, 0.000, 0.000)$ and $\text{O2}(0.500, 0.000, 0.500)$, and Au also occupies two different Wyckoff positions $\text{O1}(0.123, 0.000, 0.559)$ at 100 GPa. In the $\text{Pa}-3$ phase of AuO2 (figure 3(a)), the Au atom is 6-fold coordinated by O to form a distorted AuO6 octahedron. The Au–O bond length in the $\text{Pa}-3$ phase is 1.971 Å and the nearest neighbor O–O distance is 2.578 Å at 50 GPa, suggesting that there is no strong O–O interaction in $\text{Pa}-3$ AuO2 and AuO2 is not a peroxide in view of the fact that the covalent radius of O is 0.73 Å. The $\text{Pa}-3$ structure is composed of octahedral units that share common vertices, where all Au atoms are equivalent. In contrast to the $\text{Pa}-3$ phase, the $\text{C2/m}$ phase of AuO2 is a layer structure, with each layer comprising a plane of AuO4 groups and AuO6 octahedra. Two inequivalent Au and O atoms in the $\text{C2/m}$ phase (figure 3(b)) form two distinct types of coordination units. One unit consists of one Au atom and four surrounded O atoms; in another Au atom is coordinated by six O atoms, which is similar to the octahedra seen in the $\text{Pa}-3$ phase. In an earlier study of AuO it was reported that the most stable C2/c phase of AuO contains both linear AuO2 and square-planar AuO4 groups at 0 GPa, with mixed-valence Au$^{1+3+}$ as reported previously [10].

After investigating the thermodynamic stability of AuO2 and Au2O3, we checked the dynamical stability by calculating the phonon dispersion relations for these phases using the finite displacement method as implemented in the Phonopy code [27]. Our calculations confirm that there are no imaginary modes across the entire Brillouin zone, as shown in Fig. 4, demonstrating that these predicted structures are dynamically stable over the pressure range we have considered in this work. Previous studies have reported the presence of Au$^{4+}$ in AuF3, however, none were reported in AuO2$^2$ [11]. In the previous work, the suggestion has been made that Au has no single valence state but occurs with mixed +3 and +5 valence states in AuO2 under pressure. Electronic structures are critical to understanding the related physical and chemical properties of materials [28–30]. As illustrated in our PDOS at 50 and 70 GPa (figure 5), there is strong hybridization between the Au-5d and the O-2p in the valence electrons, indicating that charge is transferred from the Au-5d to O-2p orbitals in our
AuO₂ compounds. To further illustrate the detailed charge transfer between Au and O atoms, we calculated the Bader charge of the AuO₂ and Au₂O₃ phases with PBEsol functional at 50 GPa. The previous work [7] claimed that Au occurs as Au³⁺ in Au₂O₃, and we found that Au is calculated to lose ~1.32 e⁻ in our calculations. However, we further found that Au loses 1.63 e⁻ in our predicted AuO₂ phase (Pa-3). Since Au in Au₂O₃ loses ~1.32 e⁻ and corresponds to a +3 valence state, this suggests that Au in AuO₂ is likely to have +4 valence because 1.63 (AuO₂)/1.32 (Au₃⁺O₃) is almost equal to 4/3.

We have also performed the Mulliken population analysis [31] that allows to quantitatively provide information about the valence states of Au. In known Au₂O₃ (Fdd2 phase) with Au³⁺, Au loses 0.89e; by contrast, Au loses 1.15e in AuO₂ (Pa-3 phase). Their ratio (1.15/0.89) is about 4/3, indicating that the Au in AuO₂ should be Au⁴⁺. Moreover, crystal orbital Hamilton population (COHP) [32] also permits precise analysis of the strength and nature of the chemical bonding for the predicted structure, where ICOHP is the energy integral of COHP and it fundamentally representative chemical interactions between two atoms. The ICOHPs of the AuO₂ unit (in Au₂O₃ and AuO₆ unit in AuO₂) are −12.52 and −15.84 eV, respectively. The value of 15.84/12.52 is almost equal to 4/3, which suggests Au in AuO₂ is +4 valence again.

In this system, both Au₂O₃ and AuO₂ are ionic compounds. We take Au₂O₃ with known Au³⁺ as a good reference to investigate that the valence state of Au is 4+ in AuO₂. Au⁴⁺ is predicted in the Au–O system and it may improve the physical and chemical knowledge of Au. In our calculations, AuO₂ is a pyrite-like structure which has isotropic properties. Some compounds with transition metal having high oxidation state can serve as catalysts, fluorinating agents, and oxidants. Therefore, the newly found AuO₂ is very hopeful to be synthesized experimentally, which may be used as an oxidants like the other useful materials in practical application.

The PDOS and band structures at 50 and 70 GPa for the Pa-3 phase and C2/m phase are added in figures 4(a)–(d). In order to probe the electronic properties of our predicted AuO₂ phase, we calculated the PDOS and electronic band structures. We find that the low-pressure Pa-3 phase is a metal, as evidenced by the fact that there are occupied states at the Fermi level from the O-p orbital to Au-d orbital as shown in PDOS. At elevated pressures, its density of state at the Fermi level increases. In contrast, we find that the high-pressure C2/m structure is a semiconductor with a gap of 0.32–0.34 eV between 50 and 70 GPa. Band-gap opening may stem from the combined action of orbital hybridization (Au-d and O-p) and

**Figure 3.** Predicted ground-state and static high-pressure structures for AuO₂: (a) Pa-3 phase at ambient pressure and (b) C2/m phase at 100 GPa.

**Figure 4.** Phonon dispersion curves for the predicted (a) Pa-3 phase and (b) C2/m phase.
the interaction between Au and O, that completely fill the p-d hybridized valence band, similar to the finding in FeH$_6$ [33].

To further elucidate the mechanisms for the pressure-induced phase transition between the two ionic AuO$_2$ phases, we examined the evolution of the internal energy ($U$) and the product of pressure and volume ($PV$), which contribute to the enthalpy ($H = U + PV$) as a function of pressure. The $\Delta H$, $\Delta U$, and $\Delta(PV)$ are shown in figure 6, where the enthalpy of the $Pa$-3 phase is set to zero. Obviously, the phase transition from $Pa$-3 phase to $C2/m$ phase is mainly caused by the steep decrease of the $PV$ item; meanwhile, the internal energy of the $C2/m$ phase rises with increasing pressure. These results could reasonably explain that $C2/m$ phase could be stable at higher pressure.

**Conclusion**

In summary, we have identified a new structure of AuO$_2$ with $Pa$-3 symmetry using the PBEsol functional under high pressure. We find that the predicted $Pa$-3 phase of AuO$_2$ contains Au$^{4+}$, in contrast to the previous finding of mixed-valence states (+3 and +5) in the $C2/m$ phase of AuO$_2$. Due to the enhancement of hybridization between Au-d and O-p orbitals, AuO$_2$ displays an electronic transition from metal to semiconductor upon increasing pressure. Our enthalpy calculations suggest that AuO$_2$ is much more stable than a mixture of Au$_2$O$_3$ and O$_2$, indicating that Au–O compounds containing Au$^{4+}$ is hopeful to be synthesized experimentally at high pressure and high temperature from a mixture of Au$_2$O$_3$ and O$_2$.

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**References**
