

# Two gold surfaces and a cluster with remarkable reactivity for CO oxidation, a density functional theory study

## Citation for published version (APA):

Hussain, A., Muller, A. J., Nieuwenhuys, B. E., Gracia, J., & Niemantsverdriet, J. W. (2011). Two gold surfaces and a cluster with remarkable reactivity for CO oxidation, a density functional theory study. *Topics in Catalysis*, 54(5-7), 415-423. <https://doi.org/10.1007/s11244-011-9672-3>

## DOI:

[10.1007/s11244-011-9672-3](https://doi.org/10.1007/s11244-011-9672-3)

## Document status and date:

Published: 01/01/2011

## Document Version:

Publisher's PDF, also known as Version of Record (includes final page, issue and volume numbers)

## Please check the document version of this publication:

- A submitted manuscript is the version of the article upon submission and before peer-review. There can be important differences between the submitted version and the official published version of record. People interested in the research are advised to contact the author for the final version of the publication, or visit the DOI to the publisher's website.
- The final author version and the galley proof are versions of the publication after peer review.
- The final published version features the final layout of the paper including the volume, issue and page numbers.

[Link to publication](#)

## General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal.

If the publication is distributed under the terms of Article 25fa of the Dutch Copyright Act, indicated by the "Taverne" license above, please follow below link for the End User Agreement:

[www.tue.nl/taverne](http://www.tue.nl/taverne)

## Take down policy

If you believe that this document breaches copyright please contact us at:

[openaccess@tue.nl](mailto:openaccess@tue.nl)

providing details and we will investigate your claim.

# Two Gold Surfaces and a Cluster with Remarkable Reactivity for CO Oxidation, a Density Functional Theory Study

A. Hussain · A. J. Muller · B. E. Nieuwenhuys ·  
J. M. Gracia · J. W. Niemantsverdriet

Published online: 8 February 2011  
© The Author(s) 2011. This article is published with open access at Springerlink.com

**Abstract** We calculate the energetics of CO oxidation on extended surfaces of particular structures chosen to maximize their reactivity towards either O<sub>2</sub> dissociation, after which CO + O to CO<sub>2</sub> is a facile reaction, or to CO<sub>2</sub> from molecular O<sub>2</sub> and CO. We identified two configurations of Au atoms for which the energetics of these reactions are feasible. A site consisting of four Au atoms in a square geometry appears well suited for dissociating oxygen.

---

A. J. Muller is on leave from the University of Johannesburg, South Africa.

A. Hussain · J. M. Gracia  
Eindhoven University of Technology, STW 3.44, Helix  
Building, 5600 MB Eindhoven, The Netherlands  
e-mail: a.hussain@tue.nl

J. M. Gracia  
e-mail: j.m.graciabudria@tue.nl

A. J. Muller  
Department of Chemistry, University of Johannesburg  
(APK Campus), P.O. Box 524, Auckland Park,  
Johannesburg 2006, South Africa  
e-mail: mullera@uj.ac.za

B. E. Nieuwenhuys  
Eindhoven University of Technology, STW 3.50, Helix  
Building, 5600 MB Eindhoven, The Netherlands  
e-mail: nieuwe\_b@chem.leidenuniv.nl

J. W. Niemantsverdriet (✉)  
Eindhoven University of Technology, STW 3.47, Helix  
Building, 5600 MB Eindhoven, The Netherlands  
e-mail: J.W.Niemantsverdriet@tue.nl

A. Hussain · A. J. Muller · B. E. Nieuwenhuys ·  
J. M. Gracia · J. W. Niemantsverdriet  
Schuit Institute of Catalysis, Eindhoven University  
of Technology, 5600 MB Eindhoven, The Netherlands

A Au<sub>38</sub> cluster exposing this site provides the most favourable energetics for the CO oxidation.

**Keywords** O<sub>2</sub> adsorption · O<sub>2</sub> dissociation · CO oxidation · Au clusters · DFT · Gold

## 1 Introduction

Catalysts based on gold particles with a size of a few nanometers supported on metal oxides have gained enormous attention in the area of Surf. Sci. and catalysis since Haruta et al. reported their remarkably high activity for low temperature CO oxidation [1]. Since this breakthrough, the reaction has been studied extensively, on supported particles [2–5], atomically dispersed species [6, 7], and Surf. Sci. model systems [8].

Although it is generally accepted that the catalytic activity of Au depends to a large extent on the size of the nanoparticles [9], the question which property of the nanoparticles is responsible for the reactivity has not yet been answered conclusively and the issue of gold's high activity is still under debate [10]. Several explanations have been proposed, including the role of the support [11], quantum size effects, charge transfer to and from the support, support induced-strain, oxygen spill over to and from the support, the Au oxidation state [12], and the role of very low coordinated Au atoms on Au nanoparticles [13–15]. Of course, it is possible that several of the aforementioned effects occur simultaneously.

With respect to the mechanism of the CO oxidation, both atomic and molecular oxygen have been suggested as active species [5, 16, 17]. The energetically difficult step in a mechanism based on O-atoms is the dissociation of O<sub>2</sub> on the gold, as the activation energy for this step usually

exceeds the small heat of adsorption of the O<sub>2</sub>. In fact, using DFT calculations on stepped Au surfaces, Liu et al. [18] and Fajin et al. [19] have suggested that both atomic and molecular adsorbed oxygen can oxidize CO, the former having the lower activation barrier. Under experimental conditions, water and water derived species such as hydroxyls may play a role [20–22], and reducible support oxides may provide active oxygen for the reaction [23]. It seems plausible that multiple reaction channels and, therefore, multiple O<sub>2</sub> activation mechanisms may exist. O<sub>2</sub> molecules may directly dissociate on the support, leaving oxygen atoms to diffuse to the gold particles, or they may diffuse molecularly to the gold and react or dissociate at the particle-support interface, or somewhere on the gold particles. Essential intermediates may be atomic, or molecular oxygen, or some type of oxygen-containing species such as peroxides or carbonates [24, 25].

In this paper we ask the question what type of ensemble of gold atoms would be needed to either dissociate O<sub>2</sub> into O-atoms, or enable a direct reaction between O<sub>2</sub> and CO. To this end we have investigated the relevant reaction steps on various Au surfaces including low index, stepped and diatomic Au rows created on Au(100) employing periodic self-consistent DFT calculations. As some papers report that unsupported clusters may interact strongly with O<sub>2</sub> and even catalyze CO oxidation [26–29], we also include calculations on a gold cluster. We confirm that O<sub>2</sub> does not dissociate on a flat surface or even the stepped, relatively reactive (310) surface of gold that we used in previous work [30–32]. However, we find that extended gold surfaces consisting of diatomic rows of Au-atoms on a Au(100) surface are able to dissociate O<sub>2</sub> and allow CO oxidation, while a Au<sub>38</sub> cluster terminated by a similar double dimer facet as the diatomic row structure shows even more favorable energetics for the CO oxidation from CO and O<sub>2</sub>.

## 2 Computational Details

We used the Vienna ab initio simulation package (VASP) [33], which performs an iterative solution of the Kohn-Sham equations in a plane-wave basis set. Plane-waves with a kinetic energy below or equal to 400 eV were included in the calculations. The exchange-correlation energy was calculated within the generalized gradient approximation (GGA) proposed by Perdew and Wang (PW91) [34]. The electron–ion interactions for C, O and Au atoms were described by the projector-augmented wave (PAW) method developed by Blöchl [35]. This is essentially a scheme combining the accuracy of all-electron methods and the computational simplicity of the pseudo-potential approach [36].

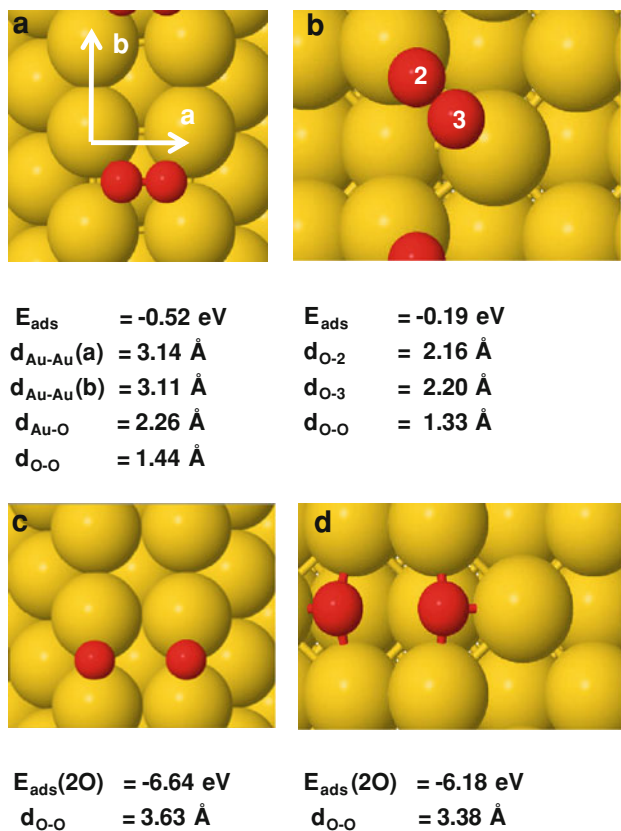
The relative positions of the Au metal atoms were initially fixed as those in the bulk, with an optimized lattice parameter of 4.18 Å (the experimental value is 4.08 Å) [37]. The optimized lattice parameter was calculated using the face-centred cubic (fcc) structure unit cell and its reciprocal space was sampled with a (15 × 15 × 15) k-point grid generated automatically using the Monkhorst–Pack method [38]. A first-order Methfessel–Paxton smearing-function with a width ≤ 0.1 eV was used to account for fractional occupancies [39]. Partial geometry optimizations were performed including the RMM-DIIS algorithm [40]. Geometry optimizations were stopped when all the forces were smaller than 0.05 eV/Å. Vibrational frequencies for transition states (TS) were calculated within the harmonic approximation. The adsorbate-surface coupling was neglected and only the Hessian matrix of the adsorbate was calculated [41]. The climbing-image nudged elastic band (cNEB) method [42] was used in this study to determine minimum-energy paths.

Molecules in gas phase were simulated in a 10 × 12 × 14 Å<sup>3</sup> orthorhombic unit cell at the Γ-point. Non-spin polarized calculations were done for closed shell CO and CO<sub>2</sub> molecules and spin-polarized calculations were performed for open shell species, O and O<sub>2</sub>.

### 2.1 Spin Polarized Calculations were Done for Adsorbates on Gold

A four layer slab model, with the two top most layers relaxed, was chosen for constructing diatomic rows on the Au(100) surface by removing one row of Au atoms along the vector b as shown in Fig. 1. A p(3 × 2) unit cell was used, and its reciprocal space was sampled with (3 × 5 × 1) k-point meshes with a vacuum gap >10 Å. The Au(100) surface was represented with a slab model using five-metal layers of which the top two were relaxed) and for Au(310) we used 11 layers with the top four relaxed, both with a vacuum gap of >10 Å to separate the periodic slabs. For the (100) slab, we used a p(2 × 2) unit cell with the reciprocal space sampled with (5 × 5 × 1) k-point meshes, and for the Au(310) surface (3 × 9 × 1) k-point meshes were used for sampling the reciprocal space of the (310) p(2 × 1) unit cell.

The Au<sub>38</sub> nano particle is a three-dimensional metal crystallite cut from metal bulk with low index surfaces (using 111 and 100 planes as basis) to have a cubo octahedral shape. The structure of the particle has been optimized in a fully relaxed state inside a 19 × 20 × 21 Å<sup>3</sup> orthorhombic unit cell, and its reciprocal space was sampled with (1 × 1 × 1) k-point mesh generated automatically using the Monkhorst–Pack method. The choice of unit cell allows for vacuum gaps of >10 Å between particles.



**Fig. 1** Top view of the two relatively reactive Au surfaces used in this paper with  $\text{O}_2$  and O adsorbed, **a**  $\text{O}_2$  adsorbed on diatomic rows on Au(100), **b**  $\text{O}_2$  on Au(310), **c** O-atoms on diatomic rows, **d** O-atoms on Au(310). Adsorption energies and characteristic distances are indicated

### 3 Results

The ability of molecular oxygen to dissociate, be it in adsorbed  $\text{O}_2$  or in a complex with CO, is crucial for  $\text{CO}_2$  formation. We will first discuss  $\text{O}_2$  adsorption and dissociation and then the reaction with CO, all on extended surfaces. Finally we discuss the reaction on the  $\text{Au}_{38}$  cluster.

#### 3.1 CO Oxidation on Au via $\text{O}_2$ Dissociation

##### 3.1.1 Adsorption of Molecular Oxygen

Figure 1 shows the two surfaces considered in this paper, namely the Au(100) with a diatomic row of Au-atoms (Fig. 1a, c) and the stepped Au(310) (Fig. 1b, d) with molecularly and dissociatively adsorbed oxygen. The interaction of molecular oxygen with gold surfaces varies markedly depending upon the surface structure, location and mode, and for many geometries the interaction is repulsive. This result is an artefact from the calculations, mainly because DFT fails to describe the attractive

long-range van der Waals interactions with respect to the short-ranged Pauli repulsion [43]. The hollow-bridge configuration of  $\text{O}_2$  on the diatomic row in Fig. 1a represents the most stable adsorption mode of  $\text{O}_2$  on these surfaces ( $-0.52 \text{ eV}$ ); single bridges as well as adsorption in double or single bridged modes are a few tenths of an eV less stable. The distances show that the Au atoms of the row relax in the direction perpendicular to the row, with Au–Au distances significantly larger than the  $2.96 \text{ \AA}$  of bulk gold. The O–O bond is clearly activated in the hollow adsorption geometry, as evidenced by the elongated O–O distance of  $1.44 \text{ \AA}$ , which is almost  $0.2 \text{ \AA}$  larger than the bond in gas phase molecular  $\text{O}_2$  (calculated at  $1.24 \text{ \AA}$ ; the experimental value is  $1.207 \text{ \AA}$  [44]). Additionally, interaction of  $\text{O}_2$  for this configuration results in a significant decrease of the stretching frequency ( $\nu_{\text{O-O}} = 748 \text{ cm}^{-1}$ ) with respect to the gas phase reference ( $1520 \text{ cm}^{-1}$  calculated vs.  $1556 \text{ cm}^{-1}$  experimental [44]) indicating substantial bond weakening.

For  $\text{O}_2$  on the Au(310) the single bridge mode in Fig. 1b is the most stable one ( $-0.19 \text{ eV}$ ), although adsorption on the long bridge at the step comes close. Both should be considered weak chemisorption modes.

Adsorption of  $\text{O}_2$  on Au(100) and on Au/Au(100), which has an additional Au atom placed in every fourth hollow site, is even weaker and is on the order of  $-0.1 \text{ eV}$ . Modes of adsorption with the O–O molecular axis normal to the surface are predicted to adsorb endothermically.

In summary,  $\text{O}_2$  interaction with gold, if it exists at all, is weak except for the diatomic row configuration presented in Fig. 1a, where  $\text{O}_2$  is chemisorbed with an adsorption energy of about  $0.52 \text{ eV}$  (about  $50 \text{ kJ/mol}$ ).

We conducted Bader charge analysis to quantify the amount of charge transfer from gold to  $\text{O}_2$  [45]. Au atoms having direct bonding with  $\text{O}_2$  on the diatomic rows are substantially more oxidized than those on the stepped (310) surface. For instance, for the most stable configuration of Fig. 1a, O-atoms are reduced by  $0.53e$  and  $0.32e$  and the Au atoms are oxidized by  $0.23e$ . However, for the most favorable configuration on (310), O-atoms gained a charge of  $0.20e$  only, with a loss of charge on the relevant Au atoms only up to  $0.13e$ . Hence we conclude that the strength of the  $\text{O}_2$  binding to Au strongly depends upon that structure of Au which is capable of donating charge to the anti-bonding  $2\pi^*$  orbitals of  $\text{O}_2$ . The charge transfer weakens the O–O bond and consequently the bond elongates.

Xu and Mavrikakis [25] have studied adsorption and dissociation of  $\text{O}_2$  on the (111) and (211) surfaces of Au (GGA = PW91). They found no adsorption on (111) but observed weak interaction ( $-0.15 \text{ eV}$ ) on the stepped surface with  $\text{O}_2$  in a top-bridge-top configuration at the step. An adsorption energy of  $-0.12 \text{ eV}$  was reported in an

earlier paper by Mavrikakis et al. [46] for  $O_2$  Au(211) with the molecular O–O axis parallel to the surface. Again, no adsorption was found on Au(111). Fajin et al. [47] have recently investigated the adsorption of  $O_2$  on Au(321) in detail (GGA = PW91). The highest  $E_{ads}$  ( $-0.17$  eV) occurred with  $O_2$  on the bridge site. A value of  $< -0.05$  eV is reported on Au(221) (GGA = PBE) [18]. A recent experimental study also reports a weak interaction of  $O_2$  with Au(111) [48]. Experimental results for the diatomic rows of Fig. 1 are not available.

### 3.1.2 Adsorption of Atomic Oxygen

Atomic oxygen, once available, adsorbs readily on the gold surfaces of Fig. 1. As we are interested in dissociation, we immediately consider the adsorption of two O-atoms on adjacent sites. Figure 1c shows the most stable configuration with two O-atoms on outer bridge positions of the diatomic row structure. The joint adsorption energy amounts to  $-6.64$  eV per two atoms (for comparison, the strongest adsorption bond for a single O-atom, which resides in the inner bridge along the b-vector amounts to  $-3.49$  eV; the adsorption energy for a single O-atom in the fourfold hollow site is  $-3.42$  eV). This configuration is exothermic by  $-0.38$  eV with respect to molecular  $O_2$  in the gas phase. The adsorption energy is  $-3.32$  eV per oxygen atom with respect to gas phase atomic oxygen. Coadsorption of two O-atoms on the fourfold hollow positions is significantly less favorable ( $-5.58$  or  $-2.79$  eV per O atom).

On Au(310) the most stable geometry of two O-atoms is as shown in Fig. 1d. A single O-atom would be adsorbed with energy of  $-3.32$  eV. However, for two adjacent O-atoms, the joint adsorption energy of  $-6.18$  eV is  $0.08$  and  $0.27$  eV endothermic with respect to  $O_2$  in the gas phase or adsorbed  $O_2$  respectively.

**Fig. 2** Potential energy diagram for CO oxidation on the diatomic Au rows on Au(100). Top (below) and side (above) views for  $O_2$  adsorption and dissociation, CO and O co-adsorption and  $CO_2$  formation are shown. The critical step is  $O_2$  dissociation which has an activation barrier almost equal to that of  $O_2$  desorption. All further steps towards  $CO_2$  however, are thermodynamically very favorable. Note that the process ends with an O-atom on the surface, making the next reaction with adsorbed CO a facile one ( $0.02$  eV activation energy and  $2.22$  eV exothermal

Adsorption energies of  $-2.47$  to  $-2.71$  eV for O on the fcc hollow site of Au(111) have been reported in literature [18, 49, 50]. Energies reported by Liu et al. [18] (GGA = PBE) on the step bridge site of Au(221) and Au(211) are  $-2.91$  and  $-3.09$  eV, respectively. Fajin et al. [47] (GGA = PW91) reported  $-3.30$  eV for O on the most stable sites of Au(321). All these values are in good agreement with our results.

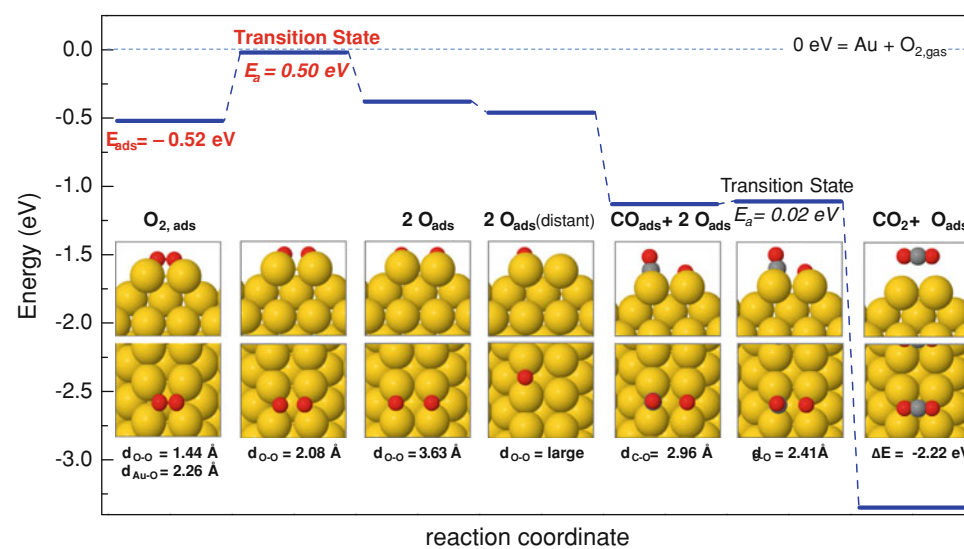
We conclude that coadsorption of two O-atoms on the diatomic rows as in Fig. 1c is expected to provide the least unfavorable thermodynamic driving force for dissociation, with an endothermicity of about  $0.14$  eV with respect to adsorbed molecular  $O_2$ , see Fig. 2.

The high adsorption energies of O-atoms on and near the steps clearly demonstrate how Au atoms of low coordination are more reactive. Again, the strongest adsorption of O-atoms is on the diatomic rows. For comparison we note however that the highest adsorption energy on this corrugated Au surface ( $-3.5$  eV) is still  $\sim 0.50$  eV lower than that on the least reactive surface of platinum, ( $4.00$ – $4.30$  eV for  $O/Pt(111)$ ), where the  $O_2$  is experimentally seen to dissociate [51].

### 3.1.3 Dissociation of Molecular Oxygen

For  $O_2$  to dissociate it is imperative that the transition state for dissociation has a lower activation energy than that for desorption of the  $O_2$ . Among the surfaces and geometries of adsorbed  $O_2$  and the two co-adsorbed O-atoms considered in this paper we found only one feasible route for dissociation, namely on the diatomic Au rows on Au(100). All other combinations led to transition states with activation energies much higher than the  $O_2$  heat of adsorption.

Figures 1a and c appear as likely start and end configurations for  $O_2$  dissociation. Both are enthalpically stable



with respect to gas phase oxygen, and the reaction is only slightly endothermic (0.14 eV). We calculate a transition state with an activation barrier of 0.5 eV, which is about the same as the activation energy for desorption of O<sub>2</sub> (0.52 eV), see Fig. 2.

In the transition state, which is symmetric, the O–O bond length increases from 1.44 to 2.08 Å and each O-atom is at a distance of 2.07 Å from the nearest Au atoms. To ascertain whether the above arrangement is a saddle point or a local minimum, vibrational analysis was carried out. The transition state was characterized by a unique imaginary frequency of 265*i* cm<sup>-1</sup> and the normal mode analysis shows that the O-atoms move in opposite directions. As the activation energy for dissociation is almost similar to that for desorption, both processes will compete. In general, desorption has a higher pre-exponential factor than surface dissociation, caused by the entropy difference between adsorbed and free molecules. Nevertheless, it is well possible that part of the O<sub>2</sub> will dissociate into O-atoms on the surface.

### 3.1.4 CO Adsorption

CO adsorbs weakly on close-packed surfaces of gold, but when the surface contains steps, adsorption energies become appreciable. On Au(310) CO bound linearly through the C-atom to the low-coordinated Au atoms at the step yields the highest adsorption energy, namely -0.73 eV. For details we refer to our previous work [32].

On the diatomic rows CO binds preferentially on bridge sites, with an adsorption energy of -0.75 eV. The bridge configuration is a local minimum with real frequencies, the CO stretch being at 1890 cm<sup>-1</sup>. The values of  $E_{\text{ads}}$  are on the order of 0.2 eV higher than on the corresponding sites of Au(100) [32]. Adsorption on the top site (-0.63 eV) has two imaginary modes at 50*i*, 95*i* cm<sup>-1</sup> and corresponds to a

second order saddle point. As the adsorption energies on bridge and top sites are very similar, we conclude that the CO molecule will be able to diffuse easily on the diatomic rows.

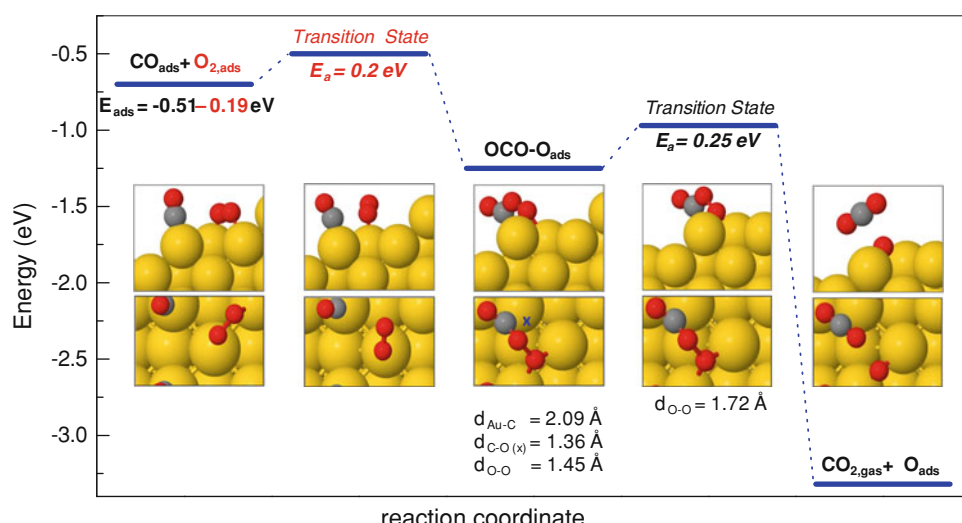
### 3.1.5 CO Oxidation on Diatomic Rows

The diatomic row may catalyze the CO oxidation reaction as illustrated in Fig. 2. Once CO and O-atoms are coadsorbed on the diatomic row, reaction takes place spontaneously with a negligible activation barrier of 0.02 eV. Similarly low activation energies have been observed in other studies [19, 52]. In the most favorable coadsorption state CO and O are both on bridge positions (see Fig. 2). In the transition state the carbon atom of CO and O-atom are 2.41 Å apart. It has one imaginary frequency of 112*i* cm<sup>-1</sup>, and is thus a true transition state. Two other combinations were tried, one with CO and O both on bridge locations along the vector b and the other with positioning CO on top and O on adjacent bridge. The former geometry is 0.45 eV less stable indicating substantial repulsion. The latter leads to the formation of CO<sub>2</sub> right away. CO<sub>2</sub> formation is highly exothermic (-2.22 eV) and thus thermodynamically favorable. As its adsorption energy is very small, it desorbs instantaneously upon formation.

## 3.2 CO Oxidation via OCOO Complex Formation and Decomposition

We have also investigated the direct reaction between CO and O<sub>2</sub>. Fig. 3 shows the reaction pathway on the (310) surface. A few initial configurations for carbon monoxide and molecular oxygen coadsorbed on the (310) surface were considered. Knowing that O<sub>2</sub> is adsorbed weakly on this gold surface, and that CO preferentially chemisorbs at the outer step [32], the combination of coadsorbates as well

**Fig. 3** Energy profile for CO<sub>2</sub> formation on Au(310) via a direct reaction of CO with molecular O<sub>2</sub>. The zero level corresponds to gas phase CO, O<sub>2</sub> and the clean slab



as the transition state shown at the start of the reaction in Fig. 3 come out as the most likely. The adsorption energy for CO and O<sub>2</sub> equals –0.7 eV. It is important to note that the differential adsorption energy of the O<sub>2</sub> in this configuration is –0.19 eV, which effectively corresponds to weak chemisorption. In the transition state, the carbon atom of the CO molecule interacts directly with the O–O bond of the O<sub>2</sub> molecule, with a transition state characterized by a small activation barrier of 0.20 eV and a unique imaginary frequency of 113*i* cm<sup>–1</sup>.

Hence, the transition state for the direct reaction between CO and O<sub>2</sub> has an activation energy of 0.2 eV, i.e. of the same magnitude as the energy with which O<sub>2</sub> binds to the Au (310) surface. Our results are rationalized in terms of enthalpy, and we do not consider entropy. O<sub>2</sub> molecules on the surface may react with CO, but the barrier to desorption, which is entropically favoured, is similar. Hence desorption will be more likely than reaction.

This OCOO species on the surface is 0.7 eV more stable than the co-adsorbate configuration. It has been invoked in other studies on Au(321) and Au(211) surfaces [19]. The structural parameters of the OCO–O complex are in excellent agreement with these studies. However, on Au(321) and (211) the activation energies for OCOO formation are substantially higher (0.58–0.68 eV) than the 0.20 eV on Au(310) in our work. Interestingly, an even lower barrier of 0.08 eV was calculated for OCOO formation on a Au strip supported on ZrO<sub>2</sub> [52].

Once the OCOO intermediate forms, it reacts via a 0.25 eV barrier to CO<sub>2</sub> and an adsorbed O-atom. Literature

reports barriers of 0.33–0.43 eV for dissociation of the OCOO species [19, 52]. The remaining O-atom reacts with CO almost without barrier (0.04 eV). The process is highly exothermic (–2.85 eV).

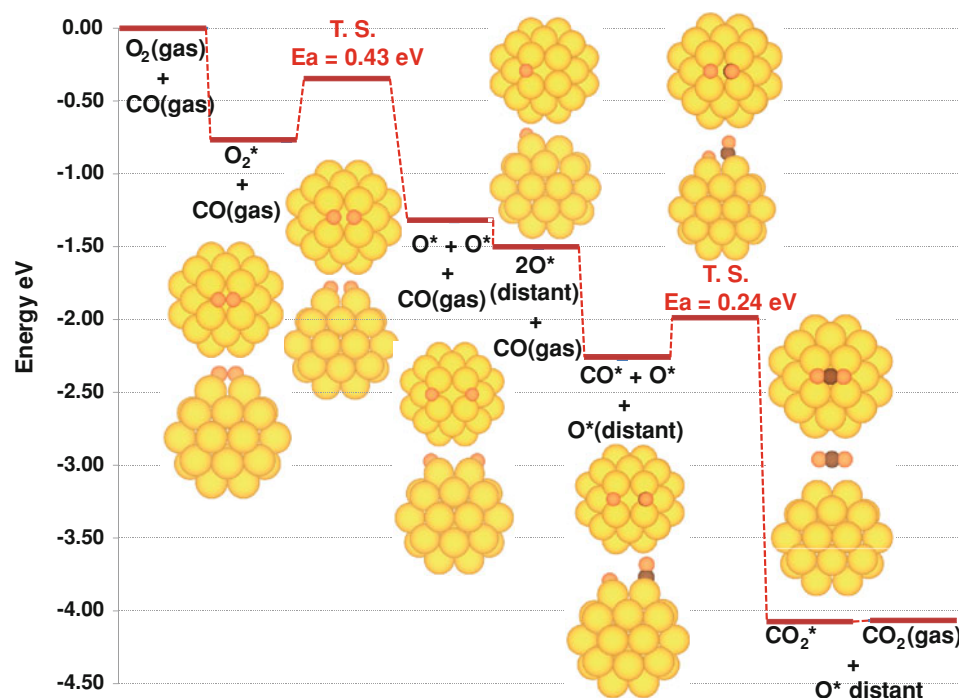
While the diatomic rows represent the most reactive Au surface for O<sub>2</sub> splitting reported thus far, this structure is almost inactive for OCOO complex formation. Thus the diatomic row structure is active enough for O<sub>2</sub> dissociation and oxidation of CO, but not via the direct reaction between O<sub>2</sub> and CO. Conversely, on the (310) surface both OCO–O compound formation and subsequent O–O bond cleavage are thermodynamically favorable and proceed by surmounting small energy barriers of 0.20 and 0.25 eV for the two steps. Hence this channel may be operative for catalyzing the CO oxidation on nanoparticles with appropriate steps.

### 3.3 CO Oxidation on a Au<sub>38</sub> Cluster

The Au<sub>38</sub> cluster shown in Fig. 4 exposes (111) facets consisting of seven atoms and (100) facets of four atoms. The latter bear similarity to the diatomic row structure on Au(100) in Figs. 1 and 2, and is therefore of particular interest for the purpose of this study.

Figure 4 starts with the adsorption and dissociation of O<sub>2</sub> on the (100) facet, in geometries that are equivalent to the ones investigated on the extended diatomic row surface. It is seen that O<sub>2</sub> adsorbs with an appreciable strength (–0.78 eV) on the cluster, while the activation energy for dissociation is only 0.43 eV, the dissociation step itself being exothermic. Hence this cluster successfully binds and

**Fig. 4** Energy scheme for CO oxidation via O<sub>2</sub> dissociation on a Au<sub>38</sub> cluster



dissociates  $O_2$  in a similar fashion as the diatomic row structure in Figs. 1 and 2. The two adjacent O-atoms after dissociation show some repulsion, hence we continue the calculations by taking twice the energy of a single O-atom per cluster. Coadsorption of CO on the same (100) facet is feasible, with a differential adsorption energy of  $-0.76$  eV. In the transition state the CO moves towards the O-atom, which involves a small barrier of  $0.24$  eV after which  $CO_2$  forms and desorbs.

We compare our results with those of Roldan et al. [29, 53], who reported  $O_2$  adsorption energies in the range of  $-0.19$  to  $-0.91$  eV on  $Au_n$  ( $n = 25, 38, 55$ , and  $79$ ) clusters.  $Au_{38}$  had the highest  $E_{ads}$  for  $O_2$  and was found to dissociate  $O_2$  with an activation barrier of  $0.46$  eV [29] in excellent agreement with our values.

Roldan et al. also investigated the effect of exchange-correlation functional on the adsorption and dissociation energy of  $O_2$  [53]. Values calculated using PW91 were larger than the PBE by roughly  $0.20$  eV, however, the effect of changing the functional on the activation barriers is smaller. Overestimation of about  $0.30$  eV in binding of  $O_2$  to gold clusters employing GGA = PW91 has been reported in previous works as well [54, 55].

#### 4 Discussion

In this paper we address CO oxidation on extended surfaces of particular structure, chosen such to maximize their reactivity towards either  $O_2$  dissociation, after which  $CO + O$  is usually a facile reaction step, or to  $CO_2$

formation between CO and molecular oxygen. We identified two configurations of Au atoms for which the energetics of these reactions appear feasible.

Following the steps of normal Langmuir–Hinshelwood mechanisms, the reactants first need to adsorb on the surface. Figure 5 presents an overview of how CO and  $O_2$  adsorb on a number of gold surfaces. Carbon monoxide chemisorbs in principle on all gold surfaces, with the exception of the close-packed  $Au(111)$  surface, where the adsorption energy is characteristic for weak chemisorption. Figure 5 clearly shows the trend that CO adsorbs more strongly when the coordination of the gold atom decreases. The strongest bond ( $-0.88$  eV) is formed with the four-fold coordination of the additional Au atom on the  $Au(100)$  surface, but adsorption of CO on the stepped (310), the diatomic row structure and the  $Au_{38}$  cluster also show considerable bond strengths between  $-0.73$  and  $-0.79$  eV. For  $O_2$  the situation is more critical. Irrespective of whether the mechanism of CO oxidation proceeds via dissociated or molecular  $O_2$ , the molecule needs to bind sufficiently strong that the activation energy of the subsequent step will not exceed the adsorption energy, otherwise the  $O_2$  will probably desorb. Of all the surfaces considered in Fig. 5, only the diatomic row and the  $Au_{38}$  cluster bind the  $O_2$  sufficiently strong. Interestingly the surface exposing the single Au atoms with the lowest coordination of all ( $N = 4$ ) does not interact appreciably with  $O_2$ . The diatomic row and the  $Au_{38}$  cluster have in common that they expose ensembles of four Au atoms of relatively low coordination in a square array to which the  $O_2$  binds in the “hollow-four centre bridge” configuration of Fig. 1a with

**Fig. 5** Overview of adsorptions and reactions relevant for CO oxidation on different gold surfaces

Au surface	Adsorption energies	CO oxidation				
		CO	$O_2$	$O$ vs $O_{gas}$	$O$ vs $O_{2,gas}$	
$Au(111)$		$-0.25$ eV non specific	0	$-2.70$ eV	$+0.43$ eV	No chemisorption of reactants
$Au(100)$		$-0.55$ eV bridged	$-0.12$ eV	$-3.14$ eV	$-0.01$ eV	CO chemisorbs, but no $O_2$ activation
$Au/Au(100)$		$-0.88$ eV on top	$-0.14$ eV	$-2.87$ eV	$+0.24$ eV	CO chemisorbs, but no $O_2$ activation
$Au(310)$		$-0.73$ eV linear, at step	$-0.19$ eV	$-3.32$ eV	$-0.19$ eV	CO + $O_2$ is energetically feasible, but as $O_2$ is physisorbed, reaction competes with diffusion and desorption of $O_2$
$Au_2/Au(100)$ diatomic rows		$-0.75$ eV Bridged	$-0.52$ eV	$-3.49$ eV	$-0.36$ eV	$O_2$ dissociation energetically feasible, but competes with $O_2$ desorption; CO + O is easy
$Au_{38}$ cluster		$-0.79$ eV bridged	$-0.78$ eV	$-3.88$ eV	$-0.76$ eV	Straightforward reaction by $O_2$ dissociation, and $CO_2$ formation

the molecular axis parallel to the surface. This site can also accommodate the O-atoms after dissociation. On the diatomic row, the four Au atoms in the square ensemble each have a coordination number of seven ( $N = 7$ ).  $O_2$  binds appreciably ( $-0.52$  eV) as do the two O-atoms after dissociation, although the dissociation is slightly endothermic ( $0.14$  eV) and the activation energy for dissociation is with  $0.5$  eV about equal to the  $O_2$  heat of adsorption. However, the same configuration on the  $Au_{38}$  cluster offers the lowest coordination of the Au atoms ( $N = 6$  for each of the four Au atoms), and here the dissociation of  $O_2$  becomes exothermic, with the concomitantly lower activation energy of  $0.43$  eV, which is considerably smaller than the  $O_2$  adsorption energy.

Our results are in good agreement with those of Roldan et al. [29, 53], who studied the  $O_2$  dissociation on a number of  $Au_n$  clusters ( $n = 25, 38, 55$ , and  $79$ ), among which  $Au_{38}$  shows the highest reactivity towards  $O_2$ . The second best is  $Au_{25}$  which interestingly also exposes the square  $Au_4$  arrangement.

Although CO oxidation via  $O_2$  dissociation on the  $Au_{38}$  cluster appears as a realistic possibility, the feasibility of this reaction on the extended diatomic rows of Au on Au (100) seems less certain. This surface does enable the adsorption and dissociation of  $O_2$  in a way that the activation energy of dissociation is similar as that for desorption of  $O_2$ . Hence, on the basis of energy one may expect that dissociation can compete with desorption. The uncertainty, however, is in the entropy change for both reactions. Desorption often has preexponential factors exceeding those of dissociation, thus favoring the former [53]. Nevertheless, we feel that our result of  $O_2$  dissociation on diatomic Au rows on Au(100) being energetically feasible is significant, as this is the only extended gold surface we have found so far which comes close to enabling the dissociation of  $O_2$ .

Another mechanism for the CO oxidation involves the direct reaction between molecular  $O_2$  and CO to  $CO_2$  and an adsorbed O-atom. The stepped Au(310) surface provides a pathway for this reaction with a barrier about equal to the  $O_2$  adsorption energy. Nevertheless, we cannot be sure that this mechanism operates, as the  $O_2$  is bound to the surface in a weakly chemisorbed state. Although reaction with CO may be energetically feasible, the molecule can diffuse freely over the surface, and hence instead of surmounting the activation barrier to form the  $OCOO$  complex, the molecule has the alternative of moving away at no energy cost.

The results in this paper highlight the importance of sites consisting of low coordinated atoms in a particular geometry, in this case a square of four Au atoms. This ensemble appears capable of binding both molecular and dissociated oxygen with sufficient strength, that

dissociation of  $O_2$  is energetically favorable. As these sites also bind CO sufficiently strong, reaction with atomic oxygen to form  $CO_2$  proceeds readily with a small activation barrier.

**Acknowledgments** We thank the National Computer Facilities NCF (Grant SH-034-08) for computer time at the Huygens Super Computer. Mr. A. Hussain acknowledges financial support from the Pakistan Higher Education Commission (HEC) and Dr A.J. Muller acknowledges financial support from Sasol Technology R&D, South Africa to enable their stay at the Eindhoven University of Technology.

**Open Access** This article is distributed under the terms of the Creative Commons Attribution Noncommercial License which permits any noncommercial use, distribution, and reproduction in any medium, provided the original author(s) and source are credited.

## References

1. Haruta M, Kobayashi T, Sano H, Yamada N (1987) Chem Lett 16:405
2. Bond GC, Louis C, Thompson DT (2006) *Catalysis by gold*, Sci., Series vol 6. Imperial College Press, London
3. Grisel RJH, Nieuwenhuys BE (2001) J Catal 199:48
4. Fu Q, Saltsburg H, Flytzani-Stephanopoulos M (2003) Science 301:935
5. Bocciuzzi F, Chiorino A, Manzoli M, Lu P, Akita T, Ichikawa S, Haruta M (2001) J Catal 202:256
6. Fierro-Gonzalez JC, Anderson BG, Ramesh K, Vinod CP, Niemantsverdriet JW, Gates BC (2005) Catal Lett 101:265
7. Guzman J, Anderson BG, Vinod CP, Ramesh K, Niemantsverdriet JW, Gates BC (2005) Langmuir 21:3675
8. Valden M, Lai X, Goodman DW (1998) Science 281:1647
9. Chen M, Cai Y, Yan Z, Goodman DW (2006) J Am Chem Soc 128:6341
10. Kung MC, Davis RJ, Kung HH (2007) J Phys Chem C 111:11767
11. Mihaylov M, Ivanova E, Hao Y, Hadjivanov K, Knozinger H, Gates BC (2008) J Phys Chem C 112:18973
12. Weiher N, Beesley AM, Tsapatsaris N, Delannoy L, Louis C, van Bokhoven JA, Schroeder SLM (2007) J Am Chem Soc 129:2240–2241
13. Lopez N, Janssens TVW, Clausen BS, Xu Y, Mavrikakis M, Bligaard T, Norskov JK (2004) J Catal 223:232
14. Hvolbaek B, Janssens TVW, Clausen BS, Falsig H, Christensen CH, Norskov JK (2007) Nano Today 2:14
15. van Bokhoven JA (2009) Chimia 63:257
16. Deng XY, Min BK, Guloy A, Friend CM (2005) J Am Chem Soc 127:9267
17. Kim TS, Stiehl JD, Reeves CT, Meyer RJ, Mullins CB (2003) J Am Chem Soc 125:2018
18. Liu ZP, Hu P, Alavi A (2002) J Am Chem Soc 124:14770
19. Fajin JLC, Cordeiro MNDS, Gomes JRB (2008) J Phys Chem C 112:17291
20. Date M, Haruta M (2001) J Catal 201:221
21. Date M, Okumura M, Tsubota S, Haruta M (2004) Angew Chem Int Ed 43:2129
22. Huang JH, Akita T, Faye J, Fujitani T, Takei T, Haruta M (2009) Angew Chem Int Ed 48:7862
23. Haruta M, Yamada N, Kobayashi T, Iijima S (1989) J Catal 115:301

24. Liu LM, McAllister B, Ye HQ, Hu P (2006) *J Am Chem Soc* 128:4017
25. Xu Y, Mavrikakis M (2003) *J Phys Chem B* 107:9298
26. Mills G, Gordon MS, Metiu H (2002) *Chem Phys Lett* 359:493
27. Quintet E, Piccolo L, Daly H, Meunier FC, Morfin F, Valcarcel A, Diehl F, Avenier P, Caps V, Rousset JL (2008) *Catal Today* 138:43
28. Xu C, Xu X, Su J, Ding Y (2007) *J Catal* 252:243
29. Roldan A, Gonzalez S, Ricart JM, Illas F (2009) *Chemphyschem* 10:348
30. Vinod CP, Niemantsverdriet JW, Nieuwenhuys BE (2005) *Phys Chem Chem Phys* 7:1824
31. Vinod CP, Hans JWN, Nieuwenhuys BE (2005) *Appl Catal A-Gen* 291:93
32. Hussain A, Curulla Ferre D, Gracia J, Nieuwenhuys BE, Niemantsverdriet JW (2009) *Surf Sci* 603:2734
33. Kresse G, Hafner J (1993) *Phys Rev B* 47:558
34. Perdew JP, Wang Y (1992) *Phys Rev B* 45:13244
35. Blochl PE (1994) *Phys Rev B* 50:17953
36. Kresse G, Joubert D (1999) *Phys Rev B* 59:1758
37. Structure data of elements and intermetallic phases, Landolt-Börnstein, New Series, Group B, Vol. III, Springer-Verlag, Berlin, 1971
38. Monkhorst HJ, Pack JD (1976) *Phys Rev B* 13:5188
39. Methfessel M, Paxton AT (1989) *Phys Rev B* 40:3616
40. Pulay P (1980) *Chem Phys Lett* 73:393
41. Head JD (1997) *Int J Quant Chem* 65:827
42. Henkelman G, Uberuaga BP, Jonsson H (2000) *J Chem Phys* 113:9901
43. Silvestrelli PL, Benyahia K, Grubisic S, Ancilotto F, Toigo F (2009) *J Chem Phys* 130:074702
44. Huber KP, Herzberg G (1979) Molecular spectra and molecular structure IV. Constants of diatomic molecules. Van Nostrand Reinhold, New York
45. Bader RFW (1990) Atoms in molecules: a quantum theory. Oxford University Press, Oxford
46. Mavrikakis M, Stoltze P, Norskov JK (2000) *Catal Lett* 64:101
47. Fajin JLC, Cordeiro MNDS, Gomes JRB (2007) *J Phys Chem C* 111:17311
48. Gong JL, Mullins CB (2009) *Acc Chem Res* 42:1063
49. Kandoi S, Gokhale AA, Grabow LC, Dumesic JA, Mavrikakis M (2004) *Catal Lett* 93:93
50. Wang GC, Tao SX, Bu XH (2006) *J Catal* 244:10
51. Getman RB, Schneider WF, Smeltz AD, Delgass WN, Ribeiro FH (2009) *Phys Rev Lett* 102:076101
52. Wang CM, Fan KN, Liu ZP (2007) *J Am Chem Soc* 129:2642
53. Roldan A, Ricart JM, Illas F (2009) *Theor Chem Acc* 123:119
54. Wallace WT, Leavitt AJ, Whetten RL (2003) *Chem Phys Lett* 368:774
55. Ding XL, Li ZY, Yang JL, Hou JG, Zhu QS (2004) *J Chem Phys* 120:9594