Surface Chemistry of Catalysis by Gold*

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1. Introduction

Gold has long been regarded as an "inert" surface and bulk gold surfaces do not chemisorb many molecules easily. However, in the last decade, largely through the efforts of Masatake Haruta, gold particles, particularly those below 5 nm in size, have begun to garner attention for unique catalytic properties (1-8). In recent years, supported gold particles have been shown to be effective as catalysts for low temperature CO oxidation (9), selective oxidation of propene to propene oxide (10), water gas shift (11), NO reduction (12), selective hydrogenation of acetylene (or butadiene) (13) and other reactions (1-5,14-16). Currently used in Japanese toilets for odour reduction (3), gold has demonstrated industrial potential as well for the hydrochlorination of ethyne to vinyl chloride (16-18) and as a bimetallic component of vinyl acetate monomer production catalysts (19,20). Low temperature CO oxidation is of particular importance, finding applications in indoor air quality applications (21) and as a guard bed catalyst to prevent CO poisoning of proton exchange membrane fuel cells (22-24).

Numerous experimental and theoretical studies are currently being performed which are aimed at understanding why gold starts to catalyse these reactions when its dimensions go to a nanometer scale. The recent surge in study of gold catalysts, which has been reviewed by Haruta (1-4), Bond (5-7) and Thompson (5,6,8), has been accompanied by a corresponding swell of interest in the surface science of gold. This review will not attempt to be completely comprehensive but rather to give the reader a view of the current state of research in the attempts to understand some of the unique catalytic chemistry of gold. Of particular note, the electrochemistry of gold and the formation of self assembled monolayers of various organics (particularly thiols) are subjects that demand their own separate reviews and the authors will not present a discussion of these areas. The review will focus on issues in heterogeneous chemistry, although gold has proven to have applications in homogeneous catalysis as well (8,16,25,26). In general, surface science studies attempt to relate properties of welldefined model structures to more complex systems. Therefore both results from gold single crystals and more intricate systems involving deposited gold particles will be presented with a discussion of their relevance to real catalysts.

Although the catalytic nature of gold may be linked to size and structure issues, it seems appropriate to begin any discussion with some general observations about gold's unique chemistry. Due to the large size of its nucleus, relativistic effects become critically important to an evaluation of gold chemistry (27-30). In short, the s orbitals and to a lesser extent the p orbitals are found to contract in response to the large nuclear charge, while the d orbitals (and f orbitals) are in fact expanded. In the case for gold, the energy difference between the 6s and 5d orbitals shrinks by over 60% (28) when compared to the theoretically modeled non-relativistic case. Calculations reveal that these effects experience a local maximum for gold when examining its place in the Periodic Table (29). Consequently, the 5d band becomes a focal point of gold chemistry and gold displays many extraordinary characteristics. For example, the higher cohesive energy of gold as compared to silver can be explained by the increased participation of the d orbitals in metal-metal bonding. Gold also has the highest electronegativity of any metal and is only slightly more electropositive than such non-metals as sulfur and iodine. Gold possesses an extremely high electron affinity (2.31 eV) and ionization potential (9.22 eV) as well (30). Gold is the only transition metal which does not have a stable oxide (Au₂O₃ is only metastable) (31). Gold also possesses a unique ability to interact with itself, known as aurophilic bonding, which allows for the formation of superstructures of gold containing molecules (32). As we will discuss later, relativistic

^{*}This review is an expansion of the Keynote lecture given by Prof. Freund at Gold 2003

effects continue to be important even when shrinking the size of gold and/or changing its chemical environment (27,33,34).

This review is organized as follows. First, we begin with the structure and adsorption properties of Au crystalline surfaces. Next, we will discuss reactions that have been observed on gold single crystals, focusing on the oxidation of CO using oxygen atoms. We will also briefly discuss the use of gold in bi-metallic systems and the contribution to this field made by surface science. Thereafter, we will examine lessons learned from gas phase experiments before moving on to theoretical predictions for the adsorption and reaction properties of gold single crystal surfaces and free gold clusters. Then, we will discuss model catalyst systems, focusing on the relationship between the support and gold with regard to structure and reactivity. Lastly, we will discuss the unresolved questions surrounding gold raised by the catalytic community and attempt to frame how surface science can help answer such questions.

Most common abbreviations used in surface science and related experimental techniques can be found in Table 1. A further point of clarification is that for the purposes of this review, "clusters" are defined to be particles with a precise number of atoms.

2 Gold single crystal chemistry

2.1 Surface structures of Au single crystals

The low Miller indexed surfaces, (111), (110) and (100), of gold, depicted in Fig. 1a-c, are all known to undergo reconstruction. Gold is the only element with a face centered cubic (fcc) crystalline structure whose (111) face reconstructs under UHV conditions (36). The reconstruction can be described by a complex stacking-fault-domain model by which both fcc and hcp domains are present on the surface with narrow transition regions in between them. The reconstructed surface, which is sometimes referred to as a $(23 \times \sqrt{3})$ structure (37), has been examined in much detail by LEED (37), helium atom scattering (38) and STM (36). The unit cell of 63 x 4.7 Å experiences contraction in the {1-10} direction, as 23 atoms sit on top of 22 hollow sites. The transition domains between regions of



Figure 1 The (a) (111), (b) (100) and (c) (110) fcc crystal surfaces. (35)

Table 1

Useful abbreviations found in the text

Abbreviation	Technique			
AES	Auger electron spectroscopy			
AFM	Atomic force microscopy			
DOS	Density of states			
DFT	Density functional theory			
DRIFTS	Diffuse reflectance Infrared transmission spectroscopy			
EELS	Electron energy loss spectroscopy			
EID	Electron induced decomposition			
EPR	Electron paramagnetic resonance			
EXAFS	Extended x-ray absorption fine structure			
FTIR	Fourier transform infrared spectroscopy			
HREELS	High resolution electron energy loss spectroscopy			
HREM	High-resolution electron microscopy			
HRSEM	High resolution scanning electron microscopy			
НОМО	Highest occupied molecular orbital			
IRAS	Infrared reflection absorption spectroscopy			
LEED	Low energy electron diffraction			
LEIS	Low energy ion spectroscopy			
LUMO	Lowest unoccupied molecular orbital			
SEM	Scanning electron microscopy			
STM	Scanning tunneling microscopy			
TEM	Transmission electron microscopy			
TPD	Temperature programmed desorption			
TPR	Temperature programmed reaction			
UHV	Ultra high vacuum			
UPS	Ultraviolet photoemission spectroscopy			
XANES	X-ray absorption near-edge structure			
XPS	X-ray photoelectron spectrocopy			
XRD	X-ray diffraction			
ML	Monolayer coverage			
L	Langmuir, 10 ⁻⁶ torr-sec			

different stacking configurations were imaged as pairs of parallel lines along the {11-2} direction. A surface corrugation of 0.2 Å was measured by STM between the high point (transition lines) and low point (fcc minima) on the reconstructed terraces. Transition lines were found to come together occasionally to form closed off U-structures with the fcc region enclosed by the U and the hcp region left outside as shown in the inset of Fig 2.



Figure 2

STM image of the reconstructed Au (111) surface (size 120 x 120 nm²). Inset (8 x 6 nm²) shows an atomic resolution image with the unit cell indicated. (36)

Large-scale images, as presented in Fig. 2, revealed a zigzag pattern (named the herringbone pattern) that results from joining of 120° rotated domains. The surface reconstruction of Au(111), while unique to fcc pure metals, is actually similar to that found when depositing a metal adlayer on top of another metal substrate with a slightly larger atomic spacing (39). The reconstruction of gold can be rationalized in terms of unique surface states that arise due to the interaction of *sp* and *d* states, a consequence of its relativistic nature (40). X-ray scattering measurements indicate that the reconstruction is stable up to 865 K, whereupon long range order is lost (41). Although the reconstruction is stable in air at room temperature (42), it can be lifted upon adsorption of certain gases. For example, exposure of above 500 mbar CO at 300 K results in an expansion of the surface layer such that it is partially returned to the unreconstructed Au(111) (43). In their examination of the stability of the reconstruction, Ertl and co-workers observed that the distance between the stacking fault lines shrank continuously with increasing potassium coverage (44). At coverages above 0.2 ML, the surface formed a new structure whereby the density of surface atoms was increased. This was attributed to an adsorbate induced weakening of bonds between the first and second layer and a simultaneous increase in bonding among surface atoms. Atomic oxygen has also been used to alter the (111) surface structure such that a $(\sqrt{3} \times \sqrt{3})R30^\circ$ structure evolves (45-49). This surface can be reduced back to the herringbone pattern described above when the sample is exposed to a reducing agent such as methanol or hydrogen (49). However, the extreme conditions under which these restructurings take place (several hours at over 800°C and 1 bar O₂) suggest



Figure 3 STM image of a "missing row" reconstruction of the Au (110)-(1x2) surface (size 15 x 15 nm²). (56)

that such surfaces are not appropriate models for catalytically active systems.

As first observed by Fedak and Gjostein (50), the Au(110) surface reconstructs into the (1x2) surface which is formed by "missing row" along {1-10} direction which is also observed for Ir(110) (51) and Pt(110) (52) and shown in Figure 3. The missing row reconstruction gives rise to surface atoms with three different coordination numbers: on top of row atoms, side of row atoms, and trench atoms. The adsorption behavior of fcc(110) surfaces is often considered in terms of different types of sites relating to the coordination number (53). The sides of row atoms give rise to a site similar to a three-fold hollow site on an fcc(111) metal and are sometimes referred to as (111) microfacets. Although this structure has been observed by LEED (54), X-ray diffraction (55) and STM (53,56), the exact details are still not entirely known. The top layer spacing is estimated to be contracted by up to 20% in the vertical direction (57). The reconstruction affects not only interatomic spacings in the surface layer but also the second layer and third layers experience compensating movements both laterally and vertically. As mentioned above, the missing row reconstruction is observed for the 5d metals but not their 4d counterparts. This suggests that the reconstruction is stabilized by the interactions involving more delocalized 5delectrons (58). The surface has been previously described to undergo an order-disorder transition at above 700 K (59). STM experiments have shown this is actually a two-step process as the reconstructed surface is stable in UHV up to ~650 K whereupon two-dimensional roughening occurs adjacent to step edges (60). Further three-dimensional roughening concomitant to surface deconstruction occurs upwards of 700 K. In an examination of CO adsorption on defects, Gottfried et al. found that defects on Au(110)-(1x2) induced by Ar⁺ ion bombardment were completely healed by heating to 500 K, thus showing the high mobility of surface atoms at relatively low temperatures (61).



Figure 4

High resolution STM image of the reconstructed Au (100)-(1x5) surface (size $5.3 \times 5.3 \text{ nm}^2$). (62)

Perhaps the most complicated of the three low indexed surfaces, the reconstruction of the (100) surface of gold is a subject of significant controversy. In Figure 4, a high resolution STM image reveals significant surface corrugation that seems to agree well with the originally suggested (5 x 1) reconstruction (50) where the added rows run along the {100} direction. Since that time, various models have been put forward, and then later a (5 x 20) (63), a (5 x 20) with rotation (64), a c(26 x 68) (37), (24 x 48) with rotation (65), and a hexagonal (28 x 5)R0.6° (66).

It has even been proposed that the reconstructed surfaces may experience different combinations of these structures depending upon step density and surface temperature (67). The reconstruction is generally believed to be limited to the first layer, thus indicating that the more compact surface arrangement is favored to a degree where the energy cost due to a lack of commensuration with the layer underneath can be overcome (68). However, Rieder et al using electron density calculations found that the best fit to their Hediffraction data occurred when they allowed for small changes in the interatomic spacing of the lower layers as well (69). The enhancement of surface bonding was seen by Annett and Inglesfield to arise as a consequence of surface band narrowing (70). Calculations by Takeuchi et al. interpreted the reconstruction primarily in terms of a relativistic effect manifested by the high degree of d orbital participation in bonding, as was the case for Au(111) (71,72). Marks and Smith proposed that the reconstruction is actually the result of surface Shockley partial dislocations (73). When the unreconstructed surface is prepared, HREM images revealed that small displacements could be seen such that one whole column of atoms appeared to be shifted both laterally and away from the surface. Upon heating to 373 K the surface transforms back to the more stable reconstructed surface (74). Marks and Smith asserted that the large degree of mass transfer that must take place for the reconstruction must occur through a mechanism involving the observed dislocations (73). X-ray scattering measurements indicated that above 1170 K the reconstruction is lifted and only the (1 x 1) phase of the bulk can be seen, thus implying the surface layer has undergone an order-disorder transition (75,76). In a similar way to the adsorption behavior of potassium on Au(111), at coverages of about 0.1 ML the reconstruction vanishes (77,78). As the coverage increases above 0.25 ML, a (1x2) reconstruction, similar to the (110) surface discussed above, forms.

2.2 Adsorption on Au single crystals

As with other metals, the adsorption of CO on gold has been extensively studied. Early work of Trapnell on gold films showed that adsorption of CO on gold is weak compared to most other metals and the heat of adsorption of CO on gold was estimated to be 8.7 kcal/mol (79). In their investigation of gold-platinum alloy films, Stephan and Ponec, observed two desorption states for CO, at 110 K and a shoulder at 170 K which they assigned to gold (80). In Bradshaw and Pritchard's examination of CO on a rough gold film, they also observed a slight red shift in the IR spectra with increasing coverage (81). An adsorption isotherm at 113 K revealed linear behavior, indicating that despite the film's roughness, CO appeared to adsorb on only a single type of adsorption site. Dumas et al (82) examined CO adsorption on rough and annealed gold films. They found that while on smooth films no CO adsorption was observable by IR spectroscopy even at 25 K, on rough films an adsorption state existed at 2125 cm⁻¹ up to 170 K. In contrast to the adsorption behavior of CO on many other metals, a small shift toward lower wave numbers



Figure 5

(a) Temperature programmed desorption spectra following CO exposure of Au(332) at 92 K as a function of CO coverage.
(b) FTIR spectra of CO adsorption on Au(332) as a function of coverage.
(83,84)

with increasing CO coverage was observed. In Ruggerio and Hollins' work on Au(332) (83,84), 3 species were identified by reflection absorption infrared spectroscopy (RAIRS), including a central peak that shifted from 2124 cm⁻¹ to 2110 cm⁻¹ as coverage increases, with both a red shifted shoulder and a blue shifted shoulder as shown in Fig 5b. Interestingly, the corresponding TPD spectra shown in Fig 5a revealed only two main peaks, one at about 140 K, the second at 185 K.

More recently, Gottfried et al. have examined the adsorption of CO on Au(110)-(1x2) in great detail (61). Five separate adsorption states were identified: α (multilayer at 32 K), β (2nd layer at 37 K), δ (physisorbed first layer at 55 K), σ (also physisorbed first layer at 67 K), and ϵ (chemisorbed at 145 K). No ordered overlayer structures were detected with LEED for any coverage examined. CO did not lift the (1 x 2) reconstruction as has been seen in the case of CO on Pt (110) (85,86). The ϵ peak was found to saturate at about 0.13 ML, a much lower saturation coverage when compared to CO adsorption on other metals. As Outka and Madix failed to detect CO adsorption at temperatures down to 125 K on Au(110), one could conclude that the ϵ feature is associated with defects (87). However, after ion bombardment, a new peak formed at ~ 190 K with an additional shoulder extending out to 230 K. In addition, ion bombardment did not lead to growth of the ϵ peak, thus indicating that the ϵ peak is not related to defects, but rather comes from a regular feature of the Au(110)-(1x2) surface. Polarization resolved UPS measurements showed that chemisorbed CO may prefer an orientation parallel to the surface. Further support of a parallel orientation was given by a lack of intensity for the 5σ orbital contribution, based upon assignments from the measurement of physisorbed CO. CO was conceived to bond to the surface primarily through back donation from Au 5*d* levels into the CO 2 π^* orbital. Previously, UPS experiments had revealed that although the surface was capable of stronger CO chemisorption on Au(110), the saturation chemisorbed coverage on Au(110) was significantly lower (only 0.3 ML) (88,89).

Goodman's lab determined a slightly lower adsorption energy than that found by Gottfried *et al* for the low coverage limit on Au(110)-(1x2) (90) (10.9 kcal/mol as compared to ~ 14 kcal/mol). Similar to Bradshaw and Pritchard (81) and Hollins and Ruggerio (83,84), a red shift in the CO stretching frequency was observed with increasing coverage. This phenomena was interpreted as the combined result of a strong "wall effect" arising from steric repulsion between the CO 5 σ and the filled *d* band of gold, a weak 5 σ donation from CO to gold and a significant 2π backdonation from the metal to CO.

CO adsorption has also been examined on Au(100) (91). By measuring the surface potential in a CO environment over a range of temperatures, the heat of adsorption was estimated to be 58 kJ/mol. This value is in agreement with findings by Gottfried *et al* (59 kJ/mol) for CO on Au(110)-(1x2) (61). As was the case for Au(110), no ordered LEED structure was observable for CO on Au(100).

At elevated pressure CO can interact in a stronger manner with gold surfaces. Peters *et al* found by surface X-ray diffraction evidence for lattice expansion of the Au(111) surface exposed to CO pressures between 0.1-530 mbar at 300 K (43), although the long range herringbone pattern was preserved. More extensive reconstruction of the Au(111) surface was observed when exposed to 110 mbar CO at 325 °C. Since the surface remained disordered after evacuation, the authors concluded that dissociative adsorption must have occurred.

Jugnet *et al.* observed similar behavior for Au(110) for CO pressures above 0.1 mbar (92). STM results showed significant surface roughening and a lifting of terrace anisotropy. This new roughened surface was stable following removal of CO. Furthermore, using polarized IRAS, a single CO adsorption state at 2110 cm⁻¹ was seen, which the authors tentatively assign to on-top adsorption as seen for other fcc(111) surfaces (93,94). However, as exposure time increased, the signal decreased, indicating possible dissociation of CO on the surface. STM images confirmed that the surface reorganization was accompanied by surface deposits which were believed to be carbon.

Much work has been devoted to exploring the nature of oxygen on gold surfaces in order to better understand the unique oxidation catalysis of gold nanoparticles. However, the surface of gold single crystals cannot be populated with chemisorbed molecular or dissociated oxygen, no matter how large the exposure to molecular oxygen (79,95), except at very high temperatures as described in Comsa's work (45-49). Although some early research reported chemisorption of oxygen (96-98), later work has ascribed these results to oxygen dissociation on calcium or silicon impurities (99,100). This has led researchers to employ other means to deliver oxygen to the surface. Canning et al. used a hot filament to crack oxygen and thereby populated a bulk gold sample with atomic oxygen (95). The surface oxide could by decomposed completely by heating and oxygen was observed to desorb associatively in a single desorption peak at 650 K. Similar results were later obtained in the same way for atomic oxygen on Au(110), yielding a single peak at 590 K (87).

Koel and co-workers have used ozone to deliver oxygen to Au(111) (101-103). They observed the formation of a single peak at 550 K that shifted to slightly lower temperatures for very low coverages. No ordered overlayer for oxygen on Au(111) was observed by low energy electron diffraction (LEED), and an oxygen coverage as small as 0.1 ML was sufficient to lift the surface reconstruction. Davis and Goodman later confirmed this result for the Au(111) surface and found that oxygen desorbs recombinatively in a single peak at 470 K from Au(100) (104). Again, no ordered overlayer structure was observed by LEED.

Christmann's group has induced oxygen chemisorption by irradiating a Au(110) surface covered with physisorbed oxygen with low energy electrons or UV photons at 28 K (105,106). Oxygen could then be seen to desorb in two peaks, low temperature feature at 497 K, which was assigned to decomposition of a Au₂O₃ surface oxide (which is inherently unstable) and a second peak at 553 K which is assigned to chemisorbed oxygen on the surface. Unlike earlier studies which report first order kinetics, second order kinetics were observed for oxygen desorption, which is typical for associative desorption of a diatomic. Further experiments involving the bombardment of a Au(110) surface by O^+ ions showed that four additional states of chemisorbed oxygen could be obtained at 415 K, at 545 K, at 620 K and at 680-850 K (107). The two lower temperature phases were similar, although not identical, to the surface oxide and chemisorbed oxygen species seen earlier. When CO was used to titrate chemisorbed oxygen both of the higher temperature features were left behind. Moreover, the highest temperature feature was found to grow and to shift to higher desorption temperatures as the ion energy and sputtering time were increased. The authors hypothesized that this feature must be associated with deep sub-surface oxygen and that the feature exhibiting a desorption peak at ~ 620 K may be associated with oxygen released from the sub-surface by a reconstruction during the heating ramp. However, experiments revealed that the work function was only restored to the value for a clean Au(110)-(1x2) surface when all the oxygen was removed. This data seems inconsistent with a sub-surface species and led the authors to suggest that the oxygen associated with high temperature desorption may be merely a strongly bound surface species. By LEED, the authors observed that the disorder induced by the adsorbed (and possibly absorbed) oxygen from O^+ bombardment could be reestablished by heating above 650 K. With sufficient cooling, oxygen can be physisorbed on gold surfaces (106). Physisorbed oxygen on Au(110)- (1x2) could be observed to desorb in three peaks: a multilayer peak at 37 K and two peaks associated with the monolayer at 45 and 51 K. The two monolayer peaks may be due to two different types of sites on the missing row reconstruction of Au(110) such as on top of the rows and down in the trenches. No physisorbed oxygen could be transferred to a chemisorbed state by simple heating of the surface.

Bär *et al.* have observed the formation of a surface oxide of gold on a FIM tip after exposure to 1 bar of O_2 for 20 min at 450 K (108). The tip has many kink and step sites which may facilitate the dissociation process. Exposure to CO led to rapid removal of the oxide.

Like oxygen, hydrogen does not dissociate readily on Au single crystal surfaces. Sault *et al.* examined the adsorption of atomic hydrogen on Au(110) using the hot filament technique that was successful for atomic oxygen dosing (109). Hydrogen was observed to desorb in a single peak at 216 K. Stobinski found that while hydrogen and deuterium will not adsorb at all on sintered gold films, molecular chemisorption was observed on unsintered films at 78 K (110). Hydrogen and deuterium were both found to desorb between 120-125 K.

Kastanas and Koel examined the adsorption of Cl_2 on Au(111) at 120 K and at 500 K (111). They found that for a 0.25 L Cl_2 an ordered (3 x 3)R30° structure could be seen with LEED when the sample temperature was less than 230 K. Unlike the case for atomic oxygen the Au(111) spots could also be clearly seen (the reconstruction is lifted upon adsorption of chlorine). Chlorine was seen to desorb in two peaks: a molecular desorption peak at 640 K and an atomic desorption peak at 790 K. No evidence of surface chloride formation or chlorine diffusion into the crystal bulk was observed.

Hydrocarbon adsorption on gold surfaces is of significant interest due to the number of partial hydrogenation and oxidation reactions that are potentially catalyzed by gold. Davis and Goodman compared propene adsorption on Au(111) and Au(100) (104) and found that the adsorption on the clean surfaces was remarkably similar. Propene desorbed in two peaks: a multilayer at 120 K and a monolayer peak at 140-145 K. No significant dissociation of propene was found to occur on either surface. Outka and Madix studied the adsorption of ethylene and acetylene on Au(110) (112). For acetylene adsorption, a single broad desorption peak between 125-200 K was observed with no signs of decomposition products. Ethylene TPD data looked nearly identical to acetylene, also with the absence of decomposition products.

Outka and Madix examined the adsorption of methanol (112), formic acid and formaldehyde (113) on Au(110) and observed reversible desorption without decomposition reactions for all of these molecules. Chesters and Somorjai also failed to observe dissociative adsorption of ethylene, cyclohexene, benzene and n-heptane on both Au(111) and the stepped surface Au(6(111) \times (100)) (96). In contrast, naphthalene was found to dissociate on both surfaces when exposed at room temperature. The resulting hydrocarbon fragments were very strongly bound indicating that the barrier to dissociation must be large.

Koel and co-workers have studied the adsorption properties of a variety of organic molecules on Au(111). Once again, acetone (114), benzene (115), cyclohexane (116), cyclohexene (116), nitrobenzene (117), and nitromethane (118) have all been shown to undergo reversible adsorption without decomposition on Au(111). Koel's group also used IRAS to study the adsorption geometries of these molecules. Wang et al concluded that nitromethane must be bonded through a single oxygen to the Au(111) surface (unlike NO₂ which is doubly bonded) to the surface for all coverages considered (118). Nitrobenzene was found to exhibit a flat geometry for low coverages up to 1.0 ML, as evidenced by the lack of intensity of in-plane modes (117). At higher coverages where multilayers of condensed nitrobenzene are formed, the molecules began to exhibit a tilted geometry with regard to both the Y and Z axes. Acetone was found to bind through the oxygen atom and adopt a tilted geometry at low coverages (112). However, as coverage was increased to about 1 ML, the orientation of the molecules changed such that the C-O bond was parallel to the surface and the molecular plane is perpendicular to the surface.

Christmann's group has examined the reversible adsorption of acetonitrile and benzonitrile (119). Neither molecule was found to exhibit ordered phases or to lift the surface reconstruction. Using a combination of work function measurements and HREELS, the authors concluded that acetonitrile bonds to the surface through its nitrogen atom with the C-C-N axis likely at some inclination to the surface. Benzonitrile was found to orient parallel to the surface as it bonded through three ring carbon atoms as well as an additional overlap between the C=N and an underlying Au atom. Christmann and co-workers went on to study the reaction of these molecules with K-promoted Au surfaces as a model electrochemical system (120).

In addition to organic molecules, Koel's group has also looked at the adsorption of nitrogen oxides on Au(111). NO2 was found to adsorb reversibly exhibiting a monolayer desorption peak at 220 K and a multilayer desorption peak at 150 K (121). HREELS measurements indicated that the molecule was oriented in an upright position and bonded to the surface through its two oxygen atoms. In multilayer films significant dimerization of NO₂ to form N₂O₄ was observed. Accompanying experiments showed that NO and N₂O did not adsorb on Au(111) for temperatures above 95 K. Building on the results from Koel's lab, ESR measurements showed that although the monolayer failed to produce an ESR spectra, isolated NO2 molecules could be detected in the multilayer on Au(111) (122). In the case of a multilayer, the monolayer acts to shield the condensed molecules above it from the spin exchange between the adsorbate spin and electrons in the conduction band of gold, which results in the loss of the spectra.

The poisoning effect of sulfur on catalysts is well known (123). As with other adsorbates, however, the interaction of sulfur with gold is somewhat unique and sulfur has even been found to act as a promoter in gold based catalysts (124). Rodriguez *et al* have studied the adsorption of sulfur on

Au(111) with photoemission spectroscopy over a wide range of coverages (125). Supporting DFT calculations confirmed earlier findings by Gottschlack and Hammer (126) that sulfur preferred the three-fold hollow site at low coverages and induced a reconstruction to a $\sqrt{3} \times \sqrt{3}$ R30° pattern as seen by LEED (127). As the coverage was increased beyond ~0.35 ML, the LEED pattern vanished and a reduction in the strength of Au-S bonds, accompanied by the formation of S₂ species, was observed. At coverages above a monolayer, sulfur preferred the on-top site and sulfur oligomers began to form. Similar behavior was observed for sulfur adsorbed on polycrystalline gold.

Unlike many other metal surfaces, the adsorption of H_2S on Au(111) and Au(100) has been found to be completely reversible (128,129). On the other hand, Jaffey and Madix have observed that about 2% of a monolayer of H_2S will decompose on Au(110) (130). Hydrogen was observed to desorb at 215 K, indicating some decomposition had occurred and a H_2S desorption at 270 K was attributed to a disproportionation reaction of SH groups on the surface. Repeating the previous experiment, Dwyer and co-workers, however, did not see decomposition of H_2S on Au(110) (131) and concluded that Jaffey and Madix's results were probably due to defects. In fact in their examination of freshly sputtered surfaces, Dwyer *et al.* observed very similar adsorption behavior to Madix and Jaffey.

As thiol molecules are frequently used to build selfassembled monolayers on gold surfaces, various researchers untaken their study for sub-monolayer have characterizations. Nuzzo et al. examined the adsorption of methanethiol and dimethyl disulfide on Au(111) and found while methanethiol was observed to experience reversible adsorption, dimethyl disulfide underwent decomposition on the surface (132). Liu et al (133) found similar chemisorption features for the adsorption of methanethiol on Au(111) as Nuzzo et al. However, decomposition products of dimethyl disulfide and methane were also observed to desorb from the surface at 470 K and 540 K respectively after exposure at 100 K. The decomposition could be attributed to defects, although theoretical calculations by Sellers predicted that with the associative formation of hydrogen on the surface an essentially thermoneutral reaction resulted (134). More recent DFT calculations suggest deprotanation of methanethiol should occur on the terraces of Au(111) (135). Liu et al also found the desorption of thiophene to be influenced by the presence of defects as the main peak at 186 K had a high temperature shoulder extending out 330 K (133). The authors concluded that this latter feature must be associated with defects, although thiophene did not experience any decomposition. SO2 was found to interact only weakly with Au(111), desorbing in a monolayer peak at 142 K, only 12 K above the multilayer feature.

Table 2

Results from TPD experiments for various adsorbates on bulk gold surfaces

Adsorbate	Gold Surface	Desorption Temperature	ref	
CO	polycrystalline	170 K		
CO	Au(332)	140 K, 185 K		
CO	Au(110)	multilayer 32 K, bilayer 37 K, physisorbed monolayer 55 K, 67 K, chemisorbed 145 K, defects 190-220K		
Atomic O	polycrystalline	recombination to O ₂ @ 650 K		
Atomic O	Au (110)	recombination O ₂ @ 590 K	87	
Atomic O by e-bombardment O ₂	Au(110)	Au_2O_3 decomposition to O_2 497 K, chemisorbed O recombination to O_2 553 K	105	
Atomic O by O ⁺ sputtering	Au(110)	surface oxide 415 K, chemisorbed species 545 K, phase transformation species 620 K, bulk O (?) 850-680 K,	107	
O ₂	Au(110)	multilayer 37 K, monolayer 45 K, 51 K	106	
O ₃	Au(111)	decomposition to O atoms- 550 K as O_2	101	
Atomic O	Au (100)	O₂ 470 K	104	
CO ₂	Au (110)	105 K	144	
Atomic D	Au(110)	216 K as D ₂	109	
D ₂	Polycrystalline	125 K on defects	110	
Cl ₂	Au(111)	Cl₂ 640 K, Cl atoms 790 K	111	
H ₂ O	Au(110)	160 K	112	
C ₂ H ₄	Au(110)	125-200 K	112	
C ₂ H ₂	Au(110)	125-200 K	112	
CH₃OH	Au(110)	multilayer 175 K, monolayer 200 K	112	
CH₃OH	Au(111)	multilayer 140 K, monolayer 160 K	102	
CH ₂ O	Au(110)	160 K	113	
НСООН	Au(110)	multilaver 175 K, monolayer 210 K	113	
C3H6O	Au(111)	multilayer 132 K, bilayer 137 K, monolayer 160 K	114	
CaHe	Au(111)	multilaver 120 K. monolaver 145 K.	104	
C3H6	Au(100)	multilaver 120 K, monolaver 140 K.	104	
CaHa	Au(111)	multilaver 151 K. bilaver 155 K. monolaver 239 K	115	
C6H10	Au(111)	multilaver 143 K. monolaver 213 K.	116	
C6H12	Au(111)	multilaver143 K. monolaver 198 K.	116	
CaH14	Au(111)	multilaver 135 K, second laver 144 K, monolaver 210 K	136	
C10H8	Au(111)	decomposition to hydrocarbon fragments	96	
H ₂ S	Au(111)	multilaver 103 K. monolaver 165 K	128	
HaS	Au(110)	multilaver 130 K. monolaver 160-180 K	131	
H ₂ S	Au(110)	multilaver130 K. multiple states 155K. 180-200 K. 220 K. H ₂ 215 K. H ₂ S from reaction 270 K. S atoms	130	
H ₂ S	Au(100)	monolaver 107 K	129	
CH3SH	Au(111)	multilaver 135 K. bilaver 190 K. monolaver 228 K	132	
CH3SH	Au(111)	120 K. 139 K. 176 K. 200 K dimethyl disulfide 470 K. methane 540 K	133	
C4H4S	Au(111)	monolaver 186 K, defects 330 K	133	
C ₆ H ₅ SH	Au(110)	multilayer 190 K, multiple states 200-350 K, benzenethiol 630 K, biphenyl 615 K, dibenzothiophene 638 K, S atoms	139	
C4H9SH	Au(100)	multilayer 138 K, butene 500K, S ₂ 940 K	141	
C₂H₅SH	Au(110)	multilayer 125 K, 2nd layer 200 K, monolayer 300 K, H ₂ 195 K, H ₂ 300 K, ethane 500 K, ethylene 500 K, ethanethiol 525 K, S atoms	138	
t-butyl thiolalcohol	Au(110)	multilayer 160 K, mutiple states 180-340 K, H ₂ 230 K, H ₂ 330 K, isobutene 470 K, H ₂ S 470 K, t-butyl thiol 470 K, S atoms	140	
$C_2H_6S_2$	Au(111)	decomposition	132	
SO ₂	Au(111)	multilayer 130 K, monolayer 142 K,	133	
NO	Au(111)	< 95 K	119	
N ₂ O	Au(111)	< 95 K	119	
NO ₂	Au(111)	multilayer150 K, monolayer 220 K	119	
C ₆ H ₅ NO ₂	Au(111)	multilayer 195 K, bilayer 210 K, monolayer 290 K	117	
CH ₃ NO ₂	Au(111)	multilayer 166 K, monolayer 177 K	118	
CH3-C=N	Au(100)	multilayer 150 K, monolayer 175 K	120	
C ₆ H ₅ CN	Au(100)	multilayer 195 K, shoulder 238 K, monolayer 285 K	120	
C ₆ H₅I	Au(111)	multilayer 188 K, bilayer 212 K, monolayer 300 K, biphenyl 400 K	151	
CH ₃ I	Au(111)	multilayer 190 K, monolayer 240 K, decomposition on defects	148	
CH₃I	Au(100)	multilayer 190 K, monolayer 250 K, C ₂ H ₆ 360 K	148	
C ₂ H ₅ I	Au(111)	215 K, decomposition products mostly C ₂ H ₄ , C ₂ H ₆ 265 K	149	
C ₂ H ₅ I	Au(100)	222 K, decomposition products mostly C4H10 260 K	150	

For higher thiols evidence of surface decomposition has been seen by Nuzzo et al with AES on Au(111) (136). In their extensive study of the adsorption of sulfur containing hydrocarbons on Au(111), Lavrich et al demonstrated that the TPD of decanethiol could be substantially altered by annealing the thiol covered surface at 343 K, and thereby inducing increased cleavage of S-H bonds (137). Madix and Jaffey observed significant decomposition of ethanethiol (138), benzenethiol (139) and tert-butyl thioalcohol (140) on the more open and thereby more reactive Au(110) surface. A pattern in their reactivity emerged. In TDS spectra between 150 and 350 K, H₂ was observed to emerge from the surface as the S-H bond was broken at the surface. Then as the surface temperature was increased above 400 K, the appropriate thiolate group on the surface decomposed and a complex series of disproportionation reactions followed, yielding products that included the thiol itself as well as higher hydrocarbons and sulfided hydrocarbons. On sulfur covered surfaces, H₂S was formed in addition to hydrogen between 200 and 350 K. The thiolate was found to be stabilized to higher temperatures on sulfur covered Au(110), and the reaction products generally had shifted distributions. Unlike the rather complicated chemistry of higher thiols on Au(110), butanethiol was seen to exhibit reversible adsorption on Au(100) (141). When butanethiol was adsorbed on a sulfur pre-covered surface, decomposition was observed, resulting in the desorption of 1-butene at 500 K and the presence of sulfur on the surface, which could either desorb at 940 K or disappear into the bulk upon annealing to high temperature. Based upon the small amount of sulfur necessary to induce decomposition, it was speculated that defects or impurities could be responsible for decomposition products seen on "clean" gold single crystal surfaces.

The observation by Gellman and co-workers that kinked surfaces possessed inherent chirality (142) has begun to be exploited by researchers for the adsorption of chiral thiols. Hammer and co-workers have calculated that the S-enantiomer of 2-amino-3-dimethylphosphino-1-propanethiol adsorbs with 8.8 KJ/mol higher binding energy on Au(17 11 9) as its R-enantiomer (143). Despite its preference for bridge bonding the thiolate group is predicted to bond to a single gold atom as the bonding of the phosphino group is optimized for the S-enantiomer. On the other hand, the R-enantiomer cannot bond well to Au(17 11 9) due to a spatial mismatch between its functional groups and the active sites of the kinked surface.

In Table 1, the results for adsorption on bulk gold surfaces are summarized. Clearly very few adsorbates undergo dissociative adsorption or decomposition on bulk gold surfaces as one may expect. However, as we will discuss in the following section, bulk gold is capable of catalyzing some interesting reactions.

2.3 Reactions on Au single crystals

Under certain conditions it is possible to study at least partial reaction cycles and thereby gain insight into catalytic mechanisms using gold single crystals. CO oxidation at 300 K was observed by Parker and Koel when a Au(111) sample populated with atomic oxygen by exposure to ozone was subsequently exposed to CO (101). They saw a linear decrease in the reaction rate with oxygen coverage. Koel's



Figure 6

(a) CO₂ production during CO exposure to O/Au(110)-(1x2). CO TPD spectrum (b) and CO₂/O/Au spectrum (c) are shown for comparison.
(d) CO oxidation over O/Au(110)-(1x2) as a function of time for specified temperatures. (144,145)

group also observed a decrease in reaction rate with temperature between 250-375 K, which gave an apparent activation energy of -2.5 kcal/mol (102). This value is consistent with several reported in the literature for real catalysts and the authors concluded that this effectively rules out an Eley-Rideal mechanism since a single step reaction cannot have a negative activation energy. Of course, in this experiment oxygen has already been provided to the surface and its dissociation may be the rate limiting step in the reaction process for real catalysts.

CO oxidation has been examined on Au(110) by Madix's and later by Christmann's group. Outka and Madix observed production of CO_2 when a room temperature Au(110) single crystal covered with atomic oxygen was exposed to CO (87). Through the use of the C¹⁸O isotope, they were able to demonstrate that CO does not dissociate on the surface as $C^{18}O^{16}O$ was the only reaction product observed. Additionally, no evidence of carbonate formation, which has been proposed by some researchers as a reaction intermediate, was found. Unlike Koel's group, Outka and Madix observed a positive apparent activation energy over a temperature range from 273-440 K. Based on their estimate for the desorption temperature of CO, a real activation energy of ~ 10 kcal/mol was calculated for an oxygen coverage of 0.25 ML. However, the authors also found that the reaction rate depended not only on the instantaneous oxygen coverage but also upon the initial oxygen coverage. They concluded that this phenomenon arose from the tendency of O-atoms to form islands at high coverages and that under these conditions CO oxidation occurs via an island eating mechanism.

Gottfried et al. examined CO2 production by TPD from Au(110) crystal covered by 1.3 ML of atomic oxygen and then exposed to CO at 1x 10⁻⁶ mbar as the sample was heated (144). As shown in Figure 6a CO₂ production was observed in three peaks: 67 K, 105 K and 175 K. CO₂ from the peak at 105 K was likely desorption limited as this coincides with the normal desorption temperature of CO₂ from Au(110). Interestingly, the peak at 67 K indicated at least some CO₂ produced was sufficiently excited by the reaction to overcome the desorption barrier. The feature at 175 K was attributed to the reaction limited production of CO2 which tails off sharply due to the low concentration of CO₂ on the surface at elevated temperatures. In other experiments samples with an initial oxygen atom coverage of 0.45 ML were exposed to CO at a various temperatures and the production of CO₂ was observed over time as shown in Figure 6b (145). Gottfried et al found that the reaction rate increased with temperature until about 170 K whereupon the production of CO₂ began to decrease. At temperatures above 175 K, the surface reaction was no longer the ratelimiting step as adsorption of CO controls CO₂ production. It was also found that the presence of oxygen on the surface increased the desorption energy of CO such that the reaction rate decreased with time, not only from a depletion of oxygen, but also from a consequential decrease in the CO surface concentration (although the gas pressure was held constant). From this analysis, the authors determined an activation energy of 57 kJ/mol (13.6 kcal/mol). In addition, annealing the surface after deposition of atomic oxygen was found to decrease the reaction rate, probably due to a loss of defects on the surface which aided in the adsorption of CO.

As mentioned in Section 2.2, Bär *et al* observed reaction of CO with a surface oxide on a gold FIM tip (108). When gas mixtures of CO and O_2 were admitted to the chamber, a reaction front could be imaged on the tip. The presence of an electric field of 12-15 V/nm greatly enhanced the reaction.

Outka and Madix went on to explore the oxidation of other adsorbates as well. When acetylene was adsorbed on a partially oxygen-covered Au(110) surface, TPR experiments yielded reaction products of water and carbon dioxide, which were observed to desorb at 205 K and 525 K respectively (112). In a first step, adsorbed acetylene was dehydrogenated by oxygen to form water, which promptly desorbed. Later at much higher temperatures, the carbon left behind on the surface from dehydrogenation reacted with oxygen. In related work, no oxidation products were observed when ethylene was adsorbed on the oxygen covered surface.

The surface reaction of formic acid with oxygen atoms on Au(110) has also been studied in the Madix lab (113). Using a hot filament as before, the gold single crystal was populated with 0.25 ML O atoms, and subsequently exposed to 0.1 ML of formic acid at 100 K. The sample was then heated and oxidation of formic acid was evidenced by desorption of water at 200 K and the subsequent desorption of CO₂. H₂O and formic acid at 340 K. The evolution at 340 K of formic acid must have been reaction limited since it was so far above the desorption temperature of formic acid from the clean surface. Using isotopically labeled reactants, the authors deduced the following reactions whereby the formic acid reacts first with adsorbed oxygen to form water and a formate group. Later the formate groups could react to produce CO2 and formic acid or they could react with adsorbed oxygen to produce CO₂ and water.

- 2 HCOOH $_{a}$ + O $_{a}$ \rightarrow H₂O $_{q}$ + 2 HCOO $_{a}$
- 2 HCOO $_{a} \rightarrow CO_{2q} + HCOOH_{q}$
- 2 HCOO $_{a}$ + O $_{a}$ \rightarrow 2 CO_{2 g} + H₂O $_{g}$

Using similar conditions, formaldehyde was seen to react on the oxygen covered Au(110) surface (113). Oxidation products of H_2O at 215 K, H_2 at 230 K, and CO_2 , H_2O and HCOOH at 340 K were observed. Additionally CH_2O was seen to desorb at 300 K, most likely stabilized on the surface by adsorbed formate groups. The reaction scheme involved the reaction of formaldehyde with surface oxygen to form adsorbed formic acid and hydrogen. This formic acid could, in turn, react with surface oxygen in a similar way as above to produce formate groups which complete the scheme.

Outka and Madix also investigated the reaction of methanol with the oxygen covered surface (112). Following an exposure of methanol at 100 K to a Au(110) surface covered with 0.25 ML oxygen, water was observed to desorb at 200 K. Results using isotopes revealed that this water was produced from the hydroxyl abstraction by surface oxygen, leaving a methoxy group on the surface. Upon additional heating to 250 K, some methanol desorbed as well as methyl formate and traces of hydrogen. Interestingly, the majority of the methanol observed to desorb at this temperature was not involved in the reaction but was stabilized on the surface by the presence of other species (oxygen, methoxy). Methoxy groups were proposed to react to form formaldehyde and some additional methanol, which desorbed. Unlike on Cu(111) (146) and Ag(111) (147), formaldehyde did not desorb as a product but reacted instead with methoxy groups to form methyl formate and hydrogen or with adsorbed oxygen to form formate groups. Further heating to 340 K produced CO₂, resulting from formate decomposition or reaction of formate with oxygen (although no hydrogen was observed).

The adsorption of water on the oxygen covered surface of Au(110) was studied by Madix's group as well (112). Although no reaction occurs, the water was chemisorbed more strongly on the oxygen modified surface leading to a desorption peak at 215 K as opposed to 190 K on the clean surface. Koel and co-workers later found very similar results for the interaction of methanol, ethylene and water with atomic oxygen on the Au(111) surface as Outka and Madix's results on the Au(110) surface (102). This may indicate that these reactions studied are structure insensitive, upon adsorption of oxygen atoms.

Davis and Goodman have attempted to observe oxidation of propene on oxygen covered Au(111) and Au(100) in a similar manner (104). Davis and Goodman saw a broadening of the propene desorption peak as compared to the clean surface with an accompanying shoulder at 200 K. Furthermore, significant amounts of the reaction products H_2O , CO and CO_2 were observed to desorb. At an oxygen coverage of 0.5 ML, trace amounts of partial oxidation products such as acetone or propylene oxide were observed but not positively identified.

Other reactions can also be driven on Au single crystals. Paul and Bent examined the relationship between structure and reactivity of gold using CH₃I as a probe molecule. They found that on a well annealed Au(111) surface, CH₃I did not react (148). However, on sputtered surfaces a small amount of ethane formed at 270 K, presumably due to partial dissociation of CH₃I at defect sites and subsequent methyl coupling. On the Au(100) surface, even when well-annealed, considerable quantities of ethane are produced from CH₃I adsorption. The coupling reaction on Au(100) did not occur until 330 K, which Paul and Bent ascribed to a stronger Au-CH₃ bond on the more open Au(100) surface. They also observed that the methyl coupling reaction could be blocked by adsorption of trimethylphosphine (TMP). Heating the surface covered with CH₃I and TMP, allowed for the observation of methyl radical desorption at 415 K. Bent's group found that higher iodoalkanes were even more reactive than iodomethane on gold single crystals (149,150). For C₂-C₄ iodoalkanes both disproportionation and coupling reactions were possible and occurred in the same temperature range, although their reaction channels were not competitive. Iodoethane was shown to produce C₂H₄, C₂H₆ and C₄H₁₀ between 260-270 K on both Au(111) (149) and Au(100) (150). However, the yield of ethene and ethane was observed to saturate at a rather low coverage of C₂H₅I whereas the yield of butane continued to increase until the monolayer was saturated. On the Au(100) surface the coupling reaction was strongly favored whereas disproportionation reactions were favored on Au(111). Higher iodoalkanes also showed an increased preference for disproportionation with increasing carbon number.

In a similar fashion, Syomin and Koel examined the formation of biphenyl produced as a result of coupling reactions from fragments of iodobenzene decomposition (151). Iodobenzene was seen to partially decompose between 200 and 250 K. Since no phenyl fragments were observed by IRAS it was concluded that their coupling reaction to biphenyl must be very fast at the surface temperatures involved. Biphenyl, whose formation on the surface was observable concomitant to C₆H₅I decomposition by IRAS, was observed to desorb at 400 K. Koel's group has successfully demonstrated electron induced decomposition (EID) to be highly selective in the scission of a single C-H bond for many adsorbed hydrocarbons and thereby explored the coupling reactions of hydrocarbon groups on Au(111) (115,116,152). Electron bombardment (30 eV) of adsorbed benzene was found to yield phenyl groups as shown by IRAS (115,152). Heating the surface resulted in evolution of benzene at about 200 K, with longer exposure times yielding narrower benzene desorption peaks. Concurrently, phenyl groups were observed to undergo coupling reactions to form biphenyl which desorbed at 400 K. Through IRAS, Koel and co-workers established that the coupling reaction to biphenyl occurred at 165 K. This result was somewhat surprising since similar reactions occur at much higher temperatures on copper and silver. Syomin and Koel further examined EID of cyclohexane on Au(111) (116,152). Cyclohexane adopted a geometry parallel to the surface of gold. However, after electron bombardment, a

cyclohexyl group was formed, which orients itself such that it was bound to the surface through a single Au-C bond. Heating the surface to 160 K caused the loss of a second hydrogen such that some cyclohexene was formed, which desorbed at the normal temperature of 215 K. A second cyclohexene peak, accompanied by cyclohexane occurred at 273 K as a result of disproportionation reactions of the cyclohexyl groups. Another mass 82 peak $(C_6H_{10}^{+})$ was observed at 324 K, which was probably not due to cyclohexene, but was rather a cracking fragment of some higher unidentified hydrocarbon.

Bartram and Koel reported that NO will react with adsorbed NO₂ on Au(111) at 100 K producing N₂O₃, as seen by HREELS (119). However, heating the surface to 170 K reversed the reaction and desorption of NO was observed. Wang and Koel also examined the interaction of NO₂ with H₂O (153). After exposing an ice film on Au(111) to NO₂ at 86 K, thermal desorption spectra revealed a complicated reaction network producing, most importantly, atomic oxygen on the surface.

2.4 Bimetallic systems involving Au

The general opinion that gold is an inert material has led researchers intuitively to utilize Au to selectively block different adsorption sites and therefore change the reaction routes. Au deposited on Si(111)-(7x7) surface will block the oxidation of the neighboring silicon, even at very low coverages (154). This seems to imply that gold not only blocks adsorption of underlying atoms but also changes the chemical nature of the silicon atoms it neighbors. Gold has also been used to decorate steps of ruthenium, effectively shutting off the dissociation of nitrogen on a gold modified Ru(0001) single crystal (155). Dahl et al demonstrated that deposition of less than 0.01 ML Au on the Ru(0001) surface reduced the sticking coefficient of N₂ by over seven orders of magnitude at 500 K.

Gold doesn't always act to merely lower the surface reactivity. Surface science studies of bimetallic systems involving gold have revealed some very interesting chemistry. In their investigation of thin Pd films on Au(111), Smith and co-workers found that gold modified the CO adsorption capability of Pd (156,157). Photoemission and HREELS data indicated that CO adsorption on the Pd monolayer was considerably weaker than on bulk Pd due to a decrease of electron density near the Fermi level. In Sellidj and Koel's examination of thin Pd films grown on Au(111) (158), the binding energy of CO was found to be slightly reduced at very low Pd coverages. However, when the surface was completely covered with a Pd layer, the CO adsorption was very similar to Pd(111). The authors note that interpretation of the data is complicated by alloying processes that occur concomitant to the temperature





Figure 7

(a) STM image of a Pd(111) surface covered by 1.5 ML Au at 320 K. (b) STM image of Au/Pd(111) surface after additional deposition of 0.25 ML Au at 110 K. (Size 335 x 335 nm²). (162)

programmed desorption. Pedersen et al also observed a decrease in CO binding strength for thin platinum films on Au(111) (159). The authors explain that Pt atoms with mostly gold neighbors lack a contribution in binding energy that they would normally experience on the Pt(111) surface, which therefore results in a lowering of the CO desorption temperature. However, as coverage of platinum was increased, an increase in the binding energy of CO on a Ptmodified Au(111) surface was observed, showing a maximum CO desorption temperature at a Pt coverage of 1.3 ML about 40 K above that of Pt(111). The results were correlated with DFT calculations that indicated that the increase in CO binding energy was due to a shift in the valence states of the Pt atoms in the Pt-Au surface alloy. Pedersen *et al* believe that the *d* band shift originated from the lattice expansion by the Pt islands, which were found by STM to have a lattice constant of 2.86 Å.

In a reverse experiment, Schmitz et al. examined the adsorption of CO on very thin gold films grown on Pd(110) (160). In much the same way that palladium was less reactive when in contact with gold, TPD experiments revealed that gold is more reactive when in contact with palladium. In Ruff *et al's* characterization of Au deposits on Pd(111) (161,162) by STM, Au was found to completely cover the Pd(111) surface at coverages as low as 1.5 ML. Using EELS, the authors demonstrated that while CO did not adsorb in any significant quantity on such films, further gold deposition resulted in the creation of a unique site with a CO stretching frequency of 2120 cm⁻¹. This adsorption site was believed to be associated with gold steps or Au island edges created by second deposition step. Figure 7a and b show STM images of gold deposited at 100 K and 300 K demonstrating low and high CO adsorption capacity respectively.

The properties of thin gold films or alloys have been investigated for other systems as well. Sachtler *et al* observed that monolayer gold films on Pt(100) were about 4 times more active than the bare Pt surface for the dehydrogenation of cyclohexene to benzene (163). Okada *et al* observed that epitaxially grown thin gold films on Ir(111) could dissociate hydrogen and deuterium, in contrast with results for Au single crystals (164).

Besenbacher et al found that addition of gold to a Ni(111) single crystal resulted in a decrease in methane dissociation on the surface (165). The initial sticking probability of methane on a Ni surface with just 0.25 ML of gold fell to $\sim 10\%$ of the sticking value of the clean Ni(111) surface (166). DFT calculations revealed that the CH₄ dissociation barrier on a Ni atom increased by 16 kJ/mol when a single Au neighbor was present (167). Two neighboring gold atoms, resulted in a 38 kJ/mol increase (the barrier to dissociation on a Ni(111) atom was calculated to be ~100 kJ/mol). The calculations show that since the methane molecule interacts with the d states of nickel in the dissociation process, the loss of reactivity can be explained by a lowering of *d*-band center when Ni is in contact with Au. Furthermore, the gold modified surface also had a reduced tendency to bind carbon and form graphite (which leads ultimately to catalyst deactivation). Besenbacher et al then went on to synthesize a high surface area Au/Ni on MqAl₂O₄ catalyst (16 wt % Ni, 0.3 wt % Au) that demonstrated a remarkable improvement in durability over a pure Ni catalyst for steam reforming of methane as the reduced carbon uptake compensates for the loss of methane reactivity (165).

Baddeley *et al* found that Au deposited on Pd(111) made for a much better acetylene cyclization to benzene catalyst (168). The ideal surface was found to contain 18% gold. The presence of gold served to lower the benzene desorption temperature, which is believed to be the ratelimiting step.

3 Chemistry of gold clusters

Although we have focused our discussion to this point upon extended surfaces, any discussion of gold chemistry seems incomplete without some mention of gas phase experiments with very small gold clusters. In real catalytic systems, gold is typically supported on metal oxides such as TiO₂, Fe₂O₃, Al₂O₃ or SiO₂. However, as we will discuss later, the nature of support can dramatically influence the reactivity of gold deposits. Therefore, in order to look at intrinsic properties of gold small particles, many groups have initiated studies of unsupported gold clusters.

One of the pioneering papers on gold chemistry that effectively foreshadowed the discovery of gold's catalytic oxidation ability, was Huber *et al's* surprising finding that a single gold atom could catalyze the oxidation of CO at temperatures as low as 10 K (169). When co-condensing Au atoms in a matrix of a $CO:O_2$ 1:1 mixture, a new species was formed, a carbonylgold(II) peroxyformate, which upon heating to 30-40 K decomposed, producing two CO_2 molecules.

Since that time, other researchers have employed gas phase methods either as flow reactors or in ion traps to examine the reactivity of gold clusters. Cox et al. examined adsorption of D_2 , O_2 and CH_4 on various sized gold clusters (170,171). The authors found that for cationic gold clusters of less than 16 atoms, deuterium readily adsorbed. However, for larger clusters no reaction occurred. For neutral gold clusters only Au₃ and Au₇ clusters were reactive with D₂. Anionic clusters were found to be completely unreactive with deuterium. Anionic clusters with an even number of atoms were found to react with oxygen, although clusters with odd numbers of atoms did not adsorb oxygen. For cationic clusters only Au₁₀ adsorbed oxygen. Methane was found to react with cationic clusters in a similar way as hydrogen with saturation conditions occurring for the same number of D2 molecules as CH₄ molecules on any given cluster size. When using mixtures of CH₄ and D₂, no higher mass products than the saturation number for any given cluster were seen, thus indicating that deuterium and methane compete for adsorption. Cox et al drew an analogy between electron affinity and the reactivity of their clusters. Clusters with the highest electron affinities had the highest reactivity to deuterium and methane while clusters with the lowest electron affinities had the highest reactivity to oxygen.

Lee and Ervin later confirmed Cox *et al's* finding for the reaction of anionic gold clusters between one and seven atoms (172) with D_2 , O_2 and CH_4 . In addition, Lee and Ervin discovered that such clusters were unreactive with NH_3 and N_2 , but did react with CO. They found that Au_n^- (n=1-4) were considerably less reactive than larger clusters. The formation of dicarbonyl species for n=4,5 and tricarbonyl species for

n=6,7 were seen as well. The variations were determined to result from the particulars of the bonding. For example, CO binds to the clusters via σ donation into the cluster LUMO and π back-bonding from the *d* orbital electrons of the cluster. Both of these are weak interactions that are not considered to be highly sensitive to the structure or size of the particle. On the other hand, oxygen is believed to bond through its unpaired electron in the π^* antibonding orbital with the HOMO of the cluster. Logically, this interaction is favored when the particle possesses an odd number of electrons such that an unpaired electron lies in HOMO, available for bonding.

Lian and co-workers found that neutral gold dimers would react with CO, NH₃ and C₂H₄, but not with O₂, N₂O, N₂ or CH₄ (173). Most unusually, hydrogen was seen to react with gold dimers at a slow rate, independent of hydrogen pressure. As Ag and Cu dimers were not seen to react with hydrogen, the authors concluded that the relativistic nature of gold is likely responsible for the increased reactivity of Au₂, as the *s*-orbital is contracted and stabilized, leading to a shorter and more stable Au-H bond.

Using UPS, Smalley's group observed even-odd oscillations of up to 1.0 eV in electron affinities, with the largest oscillations seen for gold clusters with less than 20 atoms (174). This could be explained by the relativistic effects in gold, whereby strong mixing is observed between the s and d orbitals. For clusters of 20 atoms or less, d-band features are not observed. For clusters with upwards of 70 atoms the valence region was found to be essentially indistinguishable from the bulk. Smalley and co-workers observed very local maximums in the HOMO-LUMO gaps (and minimums in electron affinity) for gold clusters with closed shell sizes of 8, 20, 34 and 58 atoms. Employing this idea of shell closing to reactivity studies, Smalley and co-workers examined the adsorption of CO on cationic gold particles (175). As expected, maximums in the abundance were found for Au_7^+CO and Au_{19}^+CO .

Li *et al* recently examined 20 atom gold clusters with anion photoelectron spectroscopy (176) and found that these structures possessed a remarkably high HOMO-LUMO gap, even greater than C_{60} (Buckey Balls), which should be an indicator of exceptional stability and perhaps chemical inertness. Accompanying DFT calculations revealed a tetrahedral pyramid to be the most stable structure, although an Au₂₀ planar structure was revealed to be nearly as favorable. The authors speculated that the tetrahedral clusters may have unusual catalytic properties due to their high surface area and large number of highly uncoordinated atoms.

Employing a gas flow reactor system, Salisbury *et al* also found that adsorption of oxygen occurred readily on anionic gold clusters with an even number of atoms (with exceptions at n=16 and n=22) whereas clusters with an odd number of atoms did not react (177). In no case did the authors observe a secondary adsorption of oxygen. As discussed above, this result could be explained by a mechanism in which oxygen adsorbs molecularly as a one-electron acceptor. As the gold cluster would then have fully paired electrons in its HOMO, it is inactive to secondary adsorption. A similar examination by Stolcic et al of the interaction of anionic gold clusters of up to 27 atoms with oxygen confirmed the finding of Salisbury et al. and additionally showed that no particles with over 20 atoms adsorb oxygen (178). For these larger clusters, it was speculated that their higher electron affinity prevented oxygen adsorption, as the adsorption process is conceived to take place through a charge transfer from the gold to the anti-bonding orbital of oxygen. When the particle is so strongly electron withdrawing, no bonding can take place. Kim et al also found that the adsorption of oxygen on anionic gold clusters was strongly tied to the electron affinity (179). However, the valence band of Au_N⁻ changed completely upon adsorption of oxygen and the peak from the HOMO of the anionic cluster (particularly for n=8-18) disappeared, suggesting charge transfer from the HOMO to oxygen. At the same time, no correlation between the geometric structure of gold and its adsorption activity was found.

Wallace and Whetten studied the adsorption of CO on anionic gold clusters and found that certain preferred products such as Au₅CO, Au₁₁CO, Au₁₅CO and Au₁₅(CO)₂ were created (180,181). These results also imply that CO adsorption can be structurally sensitive which has not been previously widely recognized. Interestingly, even at very high CO exposures, complexes with high numbers of CO molecules were not formed. This seems consistent with CO adsorption on single crystal surfaces which saturate at very low coverages. Hagen et al have recently examined the adsorption of CO on anionic monomers, dimers and trimers of gold in a ion trap experiment (182). Although CO was not found to react on Au, carbonyl complexes with one or two CO molecules attached were observed for Au₂⁻ and Au₃⁻. The formation of $Au_3(CO)_2^-$ is strongly favored, which fits well with the idea of shell closing $(Au_3(CO)_2)$ has an electron count of 8). However, the authors warn that simple examination of the s orbital electron counts is insufficient as s-d hybridization in gold due to relativistic effects must not be ignored.

Taking the next logical step in an examination of CO oxidation, Wöste and co-workers recently observed that CO and O_2 could be co-adsorbed on anionic two and three atom gold clusters, although no adsorption occurred on the monomer ion (183). The reaction of these clusters with oxygen was found to be at least an order of magnitude faster than with CO and therefore no gold carbonyls were observed. After formation of $Au_3(CO)O_2$ it was possible to adsorb a second oxygen molecule to form $Au_3(CO)(O_2)_2$ indicating a

cooperative adsorption process in which the adsorbates are not competing for bonding sites. Wallace and Whetten confirmed that this result is also true for larger clusters, i.e. the presence of one adsorbate facilitates the adsorption of the other (184). However, Wallace and Whetten produced exciting evidence for CO_2 production as well. By exposing a group of Au₆⁻ clusters to CO in the upstream portion of the reactor, a distribution of Au₆(CO)_M was created. Then this group of clusters was exposed to oxygen downstream and the products revealed the presence of $Au_6(CO)_{M-1}O^{-}$, indicating (at least indirectly) an oxidation of a CO molecule and loss of CO2. An analysis of the reaction rate reveals that the reaction rate may be over 100 times that of a real gold catalyst. However, these small clusters undergo only a single reaction cycle at most so any comparison with real or model catalysts should be tempered with the knowledge that the measurement of reaction rates is usually over thousands of reaction cycles.

In a new study by Socaciu *et al* using an octopole ion trap (185), as opposed to the flow reactor system employed by Whetten and Wallace (184), the authors report the first ever measurement of turn-over-frequencies of anionic gold dimers for CO oxidation. Socaciu *et al* observed three products upon exposure of CO and O₂ to Au₂⁻ species in the trap: Au₂⁻, Au₂O₂⁻ and Au(CO)O₂⁻. A simple mechanism was proposed by the authors to explain their findings:

 $Au_{2}^{-} + O_{2} \rightarrow Au_{2}O_{2}^{-}$ $Au_{2}O_{2}^{-} + CO \rightarrow Au_{2}(CO)O_{2}^{-}$ $Au_{2}(CO)O_{2}^{-} + CO \rightarrow Au_{2}^{-} + 2 CO_{2}$

Interestingly, when the trap was cooled sufficiently (~100 K) and a high pressure of CO was employed CO could adsorb first on Au₂⁻ leading to the formation of a different type of Au₂(CO)O₂⁻ intermediate which does not react with CO to make CO₂. Unfortunately, no direct measurement of CO₂ production was possible. DFT calculations indicated a carbonate type species, as seen in Figure 8a, formed by the Eley Rideal insertion of CO into the O-O bond of Au₂O₂⁻ was the most stable type of intermediate. However, the loss of CO by the reverse reaction shown above would be unfavorable leading the authors to conclude that a gold-peroxyformate type species, as depicted in 8b, which is formed by an Eley Rideal reaction between CO and Au₂O₂⁻ without an activation barrier as seen by Ozin and coworkers (169) must also be present.

Balteanu *et al* have used Fourier Transform Ion Cyclotron Resonance Mass Spectrometry (186), to study the coadsorption of oxygen and CO. In agreement with the preceding results oxygen was observed to react much more readily with clusters that possessed preadsorbed CO, which suggested cooperative adsorption. Furthermore, as in Wallace and Whetten's work (184) the reaction of CO was found to be highly size selective with both bare anionic



Figure 8

Two forms of Au₂CO₃ relevant to CO₂ production. a) digold carbonate and b) digold-peroxyformate with calculated bond lengths indicated. (185)

clusters and with the complexes of $Au_N(CO)_M$. Although the products were efficiently co-adsorbed, no conclusive evidence for CO_2 production was observed.

The effect of moisture on the adsorption properties of gold clusters has been examined by Wallace *et al* (187). A humid source allowed for the creation of Au_NOH clusters which were then reacted with oxygen. The hydroxylated clusters exhibited the opposite trend from the bare clusters as even-N clusters were unreactive and odd-N clusters adsorbed oxygen readily. The authors propose that the high electron affinity of the OH group leads to an electron transfer such that bare odd-N Au_N clusters which had no unpaired electrons possessed an unpaired electron available for bonding after adsorption of OH.

Cationic gold clusters from 1-12 atoms have been recently examined by Sugawara *et al* for their ability to interact with H_2S (188). Again, definite patterns with regard to the ionization potential (or electron affinity) emerged. Au⁺ and Au₃⁺ were completely unreactive and, as a general rule, even numbered particles had a higher activity. Multiple reaction products were possible as more than one H_2S molecule could adsorb and in some cases release of hydrogen was seen. Sugawara *et al* also tested the reaction of these cationic clusters with H_2 . They observed hydrogenated reaction products for clusters of up to 10 atoms. In some cases multiple hydrogen adsorption was observed. For example, for five atom cationic gold clusters up to eight hydrogen atoms could be adsorbed.

4 Theory of gold surface chemistry

Theorists have also taken up the challenge to explain the unique chemistry of gold. In their examination of H_2 dissociation on Au(111), Hammer and Nørskov found that two factors were critical to the activation barrier for reaction (and the strength of adsorbate-metal interaction as well) (189). First, gold, like all coinage metals, has a filled *d*-band, which gives rise to repulsion when an adsorbate such as hydrogen comes in contact with the surface due to filling of the antibonding orbital. Second, the degree of orbital overlap is very high for gold as compared to silver or copper resulting in a large orthogonalization energy cost and thus makes the adsorption character of gold unique.

Mavrikakis et al (190) have expanded this work to examine CO, O atom and O_2 adsorption and reaction on gold surfaces. DFT calculations indicate that CO prefers to chemisorb on the stepped surface Au(211) as compared to regular (111) terraces. Calculations for CO on Au(211) indicate a bridge bonded adsorption state with a similar binding energy as what has been found experimentally by Ruggerio and Hollins on Au(332) (83). O atoms were also found to prefer the stepped surface. O2 was found to adsorb only at steps, but the oxygen is only weakly bound, implying desorption may occur before dissociation. Whatever the nature of oxygen on the surface may be, the authors concluded that steps are an important parameter in the reactivity of small gold particles and showed that the reactivity of small particles as seen by Haruta and others (191,192) could be correlated with their expected step densities as seen in Fig 9. The data correspond even better with results from experiments when one assumes that gold atoms in the first layer are unreactive (as shown by the squares in Fig 9). Nørskov and co-workers found that O adsorption on the monolayer islands is weaker than on thicker islands and CO should not adsorb at all on the monolayer islands. However, they acknowledge that if these islands are supported by a substrate which induces strain, the strained gold lattice should have an enhanced CO adsorption capability. Strain was also found to favorably effect O2 dissociation.

Mills *et al* have examined the adsorption of O_2 on Au(111) and on ad-islands on Au(111) (193). They found that surface roughness, i.e. the presence of highly uncoordinated atoms was necessary for oxygen adsorption. In accordance with previous gas phase cluster experiments, odd numbers of electrons served to increase the binding of oxygen, although adding electrons to the flat Au(111) surface had no effect on the result. Calculations showed that the reactivity of rough surfaces and clusters results from a highly localized HOMO that allows for facile charge transfer into the π^* orbital of O_2 . On smooth or extended surfaces the HOMO is delocalized such that the surface cannot bond with O_2 .



Figure 9 Calculated step density as a function of particle size. (190)

In addition to the presence of steps, Xu and Mavrikakis have demonstrated that strained surfaces should also greatly increase the binding energy of oxygen to gold as well as lower the barrier to dissociation (194). A 10% strained surface of Au(211) was found to exhibit a binding energy of about double that of its unstrained counterpart (0.26 eV to 0.15 eV) and the reaction barrier was almost halved (from 1.12 eV to 0.63 eV).

In Strømsnes *et al's* examination of oxygen adsorption on Au(100) clusters of various sizes, calculations predicted adsorption energies of ~ 60 kcal/mol for clusters with 17 or more atoms (195), in line with the experimental value estimated by Koel *et al.* (56 kcal/mol) (101). However, DFT simulations also predicted a decrease in the adsorption energy with decreasing cluster size, which is counter to observations on supported particles (196). Accompanying calculations for H-atom adsorption predict an opposite trend: decreasing adsorption energy with increasing cluster size.

Liu et al has also examined CO and O2 adsorption and reaction on gold surfaces (197). Surprisingly, they found that steps are more reactive than kinks for the adsorption of oxygen atoms, in spite of their higher coordination number. Steps were also found to possess the lowest barrier (0.93 eV) to dissociation for O₂, even lower than small Au clusters of 12 atoms. Using their calculated values, they estimated a dissociation probability of $\sim 10^{-21}$ for steps on Au(211) based on a room temperature ideal gas, indicating that for any unsupported gold surface essentially no dissociation of oxygen occurs. Like Mavrikakis et al (186), Liu et al found that CO had a strong preference for adsorption at steps, although adsorption was also strong at adatoms and kinks. Liu et al then examined the CO oxidation reaction for the Au(221) surface. In their reaction scheme involving atomic oxygen, CO and an O atom are both adsorbed in a bridge-bonded fashion along a step edge as seen in Figure 10a. In the



Figure 10

(a) Top and (b) side views of the transition state for CO_2 production from the reaction of CO with atomic oxygen on Au(221). (c-e) show the progression of reaction between CO and molecular oxygen on Au(221). See text for details. (197)

transition state CO moves to an on-top site on the step edge and oxygen moves to a bridge site between a step atom and a terrace atom. The reaction to CO₂ was found to have an activation barrier of 0.25 eV, much lower than that of Pt(111) for example (0.80 eV). In a reaction with molecular oxygen, the proposed reaction scheme is essentially an Eley-Rideal type mechanism where O_2 in the gas phase approaches CO adsorbed at a step edge and reacts leaving an oxygen atom behind on the surface as seen in Figure 10b. The activation barrier for this reaction was found to be 0.59 eV. However, when a second CO molecule is present, the barrier is lowered to 0.46 eV, showing that CO coverage could be critical to the reaction's success. It should be mentioned that the transition state proposed by the authors is not a symmetrical carbonate species, which has been proposed in other reaction schemes. Instead they find that in the transition O-O-CO complex, the two O atoms of O₂ are chemically different from the oxygen of CO.

Although such a reaction with molecular oxygen could take place on other metals, their ability to dissociate oxygen precludes it. However, since oxygen is not dissociated by gold, a reaction pathway involving molecular oxygen is conceivable. The authors concluded the adsorption of CO and possible dissociation of oxygen are not related to particle size but rather to step densities on the particles. They point out that for the real catalyst the role of the support may be to facilitate the adsorption of oxygen. In this case the reaction would proceed along the molecular oxygen pathway as described above. However, the reaction would have a Langmuir-Hinshelwood mechanism as opposed to Eley-Rideal.

Other researchers have examined the shape of free gold particles and the possible implications for adsorption behavior. Using a combination of ion mobility measurements and DFT calculations, Furche *et al* found that gold anionic

clusters of up to at least 11 atoms preferred a planar arrangement (198). The authors believe that the relativistic nature of gold is responsible for gold's late transition from two-dimensional structures to three-dimensional particles compared to other metals. Häkkinen *et al* confirmed this result with DFT calculations of their own that showed a planar or raft-like structure is the optimal geometrical arrangement for anionic gold clusters up to 13 atoms (33, 199).

For neutral clusters, Wang et al found that small clusters (up to 6 atoms) prefer a planar arrangement, while for clusters between 7 and 9 atoms 3-dimensional geometries were preferred (200). For clusters between 10 and 14 atoms a flat cage structure possesses the lowest energy configuration. For clusters with greater than 15 atoms compact nearly spherical configurations were the most stable structures. In a similar way to experiments examining ionization energies, the HOMO-LUMO gap was found to have oscillations for even and odd numbers of atoms. The sensitivity of the density of states to cluster structure was also demonstrated by Wang et al in their comparison of the DOS for icosahedral, cubooctahedral and amorphous 13 atom clusters. Häkkinen and Landman had previously calculated that neutral gold clusters of up to 7 atoms preferred a two-dimensional arrangement (201). Grönbeck and Andreoni have also confirmed a planar arrangement for both anionic and neutral gold clusters from 2 to 5 atoms (202).

An interesting question arises about the electronic nature of these two dimensional particles. Zhao *et al* carried out calculations on planar particles to determine their polarizability (203). In the lateral direction the polarizabilities were found to be large (like a metal). However, the clusters displayed non-metallic character in the normal direction. For a very large two-dimensional Au cluster, the polarizability in the normal direction is essentially equal to a semiconductor like Si. Calculated densities of states revealed discrete behavior for 4 atom clusters and, as cluster size was increased to 20 atoms, a slow transition to a band-like DOS was observed.

Theoretical predictions for the structure of larger (between 30-1000 atoms) gold clusters have also been performed. To date, there is some debate over the lowest energy arrangements of such clusters, as proposals for disordered structures (204,205) and various types of faceted particles (176,206-210) have both been put forward. As pointed out by Marks (201), the determination of the correct equilibrium structures of small particles is complicated by changes that can occur during observation due to the influence of the HREM (212) and due to the influence of the underlying support. Furthermore, there may be only small differences in energy between structures allowing for facile transformation (205). Of particular interest is the transition between fcc structures which predominant for very large clusters, and decahedral or icosahedral particles, whose formation is predicted to be favored for very small particles. Whereas Cleveland *et al.* predicted that gold should transition from fcc structures to decahedral particles for sizes below 200 atoms and to icosahedral particles at sizes below 100 atoms (210), Uppenbrink and Wales found that the transition from fcc structures to icosahedral particles should occur at 550 atoms for gold (209). As we will discuss in the following section, the details of the structure may have strong implications for its reactivity.

Computational methods have also examined adsorption on small gold clusters. DFT calculations performed by Okumura et al show that AuO_2^{-1} is considerably more stable than AuO₂ (213). Calculations of Au₁₃ clusters showed an optimized arrangement for O₂ adsorption in which the surface atoms had some partial negative charge. Based on these findings, the authors suggest that negatively charged surface atoms are active sites for oxygen adsorption for supported gold particles. In similar calculations, Mills et al. report that oxygen adsorption is strongly preferred on even numbered anionic gold clusters as opposed to odd numbered or neutral clusters as has been observed in gas phase experiments (214). Bonding is conceived to take place by means of a charge transfer from gold to the anti-bonding π^* orbital of O₂. Wells et al. calculated that while oxygen bonding was weak for nine and eleven atom anionic gold clusters, it adsorbed readily on Au10⁻ (215). Wells et al reported a different geometry for the bonding arrangement as they favor an edge on attachment to the HOMO of the cluster as opposed to the bent-end-on configuration favored by Okumura et al and Mills et al Yoon et al and Franceschetti et al have also examined the reaction of O₂ with anionic gold clusters (216,217). Calculations showed that for n>3, oxygen would prefer to be adsorbed dissociatively. However, the energy barrier for dissociation computed by Yoon et al was still quite high (2.33 eV) for $Au_6O_2^-$, which is the most favorable case (216).

Metiu and co-workers advise some caution with regard to application of DFT to questions involving the adsorption of oxygen on gold (218). DFT calculations using both B3LYP and PW91 functionals were found to lead to prediction of significant overbinding of oxygen to gold clusters as compared to experimental findings and cluster methods.

Wu *et al* considered CO adsorption in free gold clusters between one and six atoms and calculated that CO preferred to adsorb in a single-coordinated fashion regardless of cluster size or the cluster's charge state (219). The charge state did, however, influence the strength of the interaction as cationic clusters bind CO more strongly. The n_{co} vibrational frequencies were predicted to decrease from 2228 cm⁻¹ for



Figure 11

Interaction energy in eV per molecule for CO and O_2 adsorption versus coordination number for gold atoms in various geometries. (222)

n=1 to 2166 cm⁻¹ for n=6, but were always above the gas phase value (2143 cm⁻¹). The clusters themselves were found to prefer a flat or raft-like orientation, both with and without CO adsorbed.

Häkkinen and Landman have also calculated the oxidation of CO on an anionic gold dimer (220). They determined that the most favorable arrangement should be a carbonate complex $Au_2CO_3^-$. The carbonate can then react with a second CO to produce two CO_2 molecules. In alternate pathway the carbonate can simply decompose, releasing CO_2 and leaving Au_2O^- behind, which is then available for reaction with CO.

In the study of a neutral ten atom gold cluster, Lopez and Nørskov found that O₂ dissociation was very facile (221) which had not been previously seen on Au(111) or Au(211) (190,197). The increased reactivity of the ten atom cluster is attributed to a higher lying d state as compared with atoms on single crystal surfaces. However, the geometry of the cluster is highly critical to its ability to adsorb oxygen. In their model a two layer raft-like structure was chosen with three atoms lying atop a seven atom layer. This is in contrast with the Au(111) type geometry that was examined previously by Mavrikakis et al for a twelve atom cluster (190). The authors found that both O_{2} and O atoms adsorbed on Au_{10} could react with CO with reaction barriers less than 0.4 eV. Nørskov's group studied the effect of geometry in more detail, calculating the variation of the strength of adsorption with the coordination number of gold (222) as shown in Figure 11.

Examination of gold nanowires with a coordination of two, revealed that CO experienced very strong adsorption on the order of 2 eV, which is approximately equivalent to CO binding to more reactive surfaces such as nickel or platinum. Perhaps even more surprising, oxygen was found to dissociate in a highly exothermic manner and insert itself into gold chains forming an alternating string of gold and oxygen atoms. The authors suggest that the gold nanowires that have been previously cited for their long bond distances (223) may in fact be oxygen-gold chains. Recently, it has been suggested that hydrogen could also be responsible for the stabilization of large Au-Au bond distances (224).

Metiu and co-workers have interpreted their results of calculations examining the adsorption of propene on gold clusters and surfaces in a slightly different way (225). Rather than a direct relationship between coordination number and adsorption strength, Chrétien et al developed a set of semiempirical rules regarding the position of the LUMO of a particular gold cluster, as the bonding of propene was conceived to occur through donation of electron density from its HOMO into the particles LUMO. Therefore, propene adsorbed most favorably at the point at which the lowest lying LUMO extends the furthest into vacuum. Furthermore, the lower the LUMO, the higher the binding energy. Only in cases where the LUMO density would be equivalent would the coordination number become a critical factor. An adatom on a gold surface becomes a naturally favorable adsorption site as its LUMO extends into the vacuum far more than the underlying terrace, which by nature has a highly delocalized LUMO. In addition, it was found that an adatom of gold on a gold surface should be more reactive to propene than an adatom of gold above a vacancy on a TiO₂ surface as the former case is electron depleted compared to the latter and thereby accepts an electron more easily. Of course for an electron-accepting adsorbate, like O2, this would be the opposite and adsorption to Au above the defective oxide would be favored.

5 Gold model catalysts

5.1 Growth and Nucleation of Gold

Of course, in real heterogeneous catalysts, gold is usually dispersed on a metal oxide support. Therefore, questions also arise about the role of the support and the origin of any differences in catalytic activity between gold catalysts on various supports (191,226). Therefore, in addition to research on single crystal surfaces as described in Section 2, many researchers have begun to examine gold particles deposited on either metal oxide single crystals or thin films, whereby the oxide is a model for the catalytic support. Surface science studies of planar model catalysts have been demonstrated to provide a bridge of the materials gap between "real" catalysts and metal single crystals (227-235).

Of these, gold deposited on TiO_2 rutile(110) remains the most widely studied system involving gold to date. Cosandey and Madey have written an excellent review of the Au/TiO₂ system (236) and we will attempt to highlight some of this

work as well as discuss results for other model catalysts. The (110) face of rutile has been examined extensively with STM as reviewed by Diebold (237). The (1x1) surface consists of alternating rows of titanium and oxygen atoms with half of the titanium atoms covered by so-called bridging oxygen. This leads to the creation of both 6-fold coordinated Ti (as in the bulk) and fivefold coordinated Ti, as well as threefold coordinated oxygen (as in the bulk) and two fold coordinated oxygen (bridging oxygen). Many researchers believe that due to their undercoordinated bonding, the bridging oxygens can be easily removed to form point defects or color centers, which strongly influence the support's chemistry. These defects can take on different forms depending upon whether oxygen is removed as O^{-} , O^{+} , or O^{-} . Pacchioni has recently predicted that the removal of an oxygen atom on TiO₂ results in a redistribution of charge over the Ti ions surrounding the vacancy such that the binding energy of the five-fold coordinated Ti atoms experience negative binding energy shift and the six-coordinated (of which there are now fewer) undergo a positive binding energy shift (238). This is in contrast to a surface like MgO where the charge left behind is strongly localized (239). A strongly reduced TiO₂(110) surface may undergo reconstruction to the (1x2) structure, although the exact details are still an area of active research (237). Due to the strong relationship between nucleation and defect density, surface preparation and characterization are very important to chemistry observed.

Studies by Zhang et al (240), Lai et al (241), Guo et al (242), Santra et al (243), Spiridis et al (244) and Parker et al (245) all confirm that gold experiences a two dimensional growth mode or quasi-2-D growth for very low coverages (< 0.15 ML) and then subsequently changes to a 3-D growth mode. STM images appear to show growth of gold at low coverages above Ti rows (241), but no atomically resolved images of both the particle and support simultaneously have been achieved such that one can make an unambiguous assignment. Santra et al observed that the nucleation density of particles at the step edges saturated much faster than on the terraces and that the average height of particles nucleated at step edges grew much faster than particles on terraces (243). Spiridis et al observed similarly strong preferential nucleation at step edges (244). As mentioned above there is substantial evidence that F centers (3+ defects on the TiO₂ surface) act as nucleation sites for gold particles and may help stabilize the particles. Lai et al found that STM images of the TiO₂(110) surface covered with 0.1 ML of gold appeared free of defects, thus implying that the small amount of gold on the surface must have nucleated at the defects and thereby obscures them from view (241). Using low energy ion scattering, Parker et al showed that ion bombardment of a TiO₂(110) surface to produce defects led to a higher nucleation density for gold deposited on the



Figure 12

STM images of the (a) clean $TiO_2(110)$ surface and with 0.04 ML Au deposited at (b) 130, (c) 210, and (d) 300 K, showing the decrease in the number of exposed bridging oxygen vacancies (marked by squares) with increasing gold deposition temperature. (Image size $15 \times 15 \text{ nm}^2$). (246)

defective surface (245). On the clean as prepared surface, the critical gold coverage for the transition from two dimensional growth to 3-D growth was found to be 0.09 ML whereas on the sputtered surface the critical coverage was found to be 0.22 ML. Parker *et al* also saw that the temperature of the $TiO_2(110)$ crystal during gold deposition can greatly affect the nucleation density. The critical coverage for gold was found to be 0.19 ML for gold deposited at 160 K as opposed to 0.09 ML for gold deposited at 300 K, demonstrating that the growth mode is more three dimensional at higher temperatures. Indeed, Zhang *et al* reported that gold deposited at a sample temperature of 475 K will experience three-dimensional growth from the beginning on $TiO_2(110)$ (240).

Recently, Wahlström *et al* has probed the relationship between gold particle nucleation and oxygen vacancies on a $TiO_2(110)$ single crystal in detail (246). STM measurements showed the number of vacancies was drastically reduced upon the deposition of gold, even though gold occupied only a small fraction of the sample surface. As shown in Figure 12, when depositing gold at higher temperatures the reduction in vacancies was even larger, leading the authors to conclude that the particles trapped multiple oxygen vacancies beneath them. DFT calculations revealed that gold atoms prefer to bind to oxygen vacancies by more than 0.45 eV as compared to any site on the ideal titania surface. A single vacancy was found to be capable of binding up to 3 atoms. A transition between 2-D particles and 3-D particles was shown to be present for particles with less than 3 atoms per vacancy and more than three atoms per vacancy, respectively. Unlike calculations performed by Sanchez *et al* (247) for the binding of gold clusters on MgO, no significant charge transfer was seen.

The growth and nucleation behavior of gold on various supports has been modeled by numerous researchers. To date there has been a lack of agreement on the most stable adsorption sites for gold on oxide surfaces. For titania, various researchers have established that the on-top site above a fivefold coordinated Ti (248), the on-top site above a bridging oxygen (249, 250), bridge bonded between two bridging oxygen atoms (249), between a bridging oxygen and a 5 fold coordinated Ti on the basal plane (250), a four-fold hollow over 5 fold coordinated Ti and both a bridging oxygen and an in plane oxygen (251) or the pseudo hollow site generated by two in plane O-atoms and one in the bridging rows (252) have the highest adsorption energy. Lopez and Nørskov have recently found in their calculations that the preferred adsorption site may depend on the coverage of gold which may help to explain differences between various theoretical predictions (253). Overall, gold was found to show much weaker bonding to titania when compared to copper or silver regardless of the adsorption site. Furthermore, as a consequence of the weak interaction, the energy differences between various sites are often very small (250). This can be explained by gold's relatively high electronegativity and high ionization potential which make gold a poor electron donor. Wang and Hwang found that one must apply caution in prediction of nucleation sites, particularly with regard to the size of the unit cell chosen, so as to capture the interaction between Au and the substrate instead of Au-Au interactions (251).

Not surprisingly, defects were found to play an important role in the adsorption of gold on titania and magnesia. Lopez and Nørskov calculated that oxygen vacancies were about 0.5 eV more stable than any possible adsorption site on the defect-free TiO₂(110) surface (253). Lopez et al and Giordano et al (249) also found that while sites along the (001) direction of rows containing Ti atoms are essentially energetically equal, diffusion along the protruding rows must overcome a large energy barrier. This result may help to explain why gold particles on TiO₂(110) are frequently elongated in a particular direction. Rodriguez et al (252) calculated that Au atoms adsorbed on oxygen vacancies were 48 kcal/mol (2.0 eV) more stable than the best possible adsorption site for the defect free surface. Wang and Hwang demonstrated with their DFT calculations that neighboring oxygen vacancies further increased the interaction between gold and the defect site (251). Yang et al. (254), and Vittadini and Selloni (255) calculated similar stability advantages for the adsorption of gold on oxygen vacancies on MgO(001) and TiO₂ anatase (101) respectively.

In addition to their work at low gold coverages, Madey's group has also examined the growth and nucleation behavior of large gold coverages on the TiO₂(110) surface (256). High resolution SEM images revealed that above a deposition of 1.5 nm (more than 5 ML), a new stage of growth began, marked by the elongation of particles and partial coalescence resulting in worm like islands. Continued deposition of gold resulted in decreasing island density due to coalescence, until the entire network of islands was connected at about an 8 nm coverage. Continued deposition eventually results in a granular gold film at an average thickness of 12 nm. Madey and coworkers observed that upon annealing of the large worm like islands, the particle density was unchanged but the islands thickened and formed faceted structures, revealing more of the TiO2 surface underneath. This behavior is consistent with the island thickening model supported by Campbell (228) for metals grown on metal oxides. This is in contrast to the effect of annealing at lower coverages where both coalescence and ripening processes led to a decline in particle density simultaneous to the thickening process.

Although highly faceted particles have not been seen by STM for small gold particles on TiO_2 in the same way as for Pd/Al₂O₃ for example (257), HREM images of larger particles have shown that gold particles will grow in a semi-epitaxial manner (236,258,259). Particles with a Au(111) orientation have been observed to grow such that gold lies on top of bridging row oxygen atoms. Along the (001) direction of TiO₂, the lattice mismatch is only 2.6%, but in the perpendicular (110) direction the misfit between two bridging oxygen rows and four Au atoms in (110) closed packed rows is 14.3%. A more favorable arrangement with Au(112) particles growing atop TiO₂(110) has also been observed with Au atoms sitting atop 5-fold coordinated Ti atoms. The mismatch in the (001) direction is now reduced to 8.4%.

In Figure 13 the epitaxial relationships are displayed. Cosandey and co-workers found that when gold was deposited at 300 K and then annealed to 770 K the $(111)_{Au}//(110)_{TIO2}$ orientation was preferred while direct deposition at high temperature preferentially gave rise to the $(112)Au//(110)_{TIO2}$ orientation (259). Interestingly, the Au lattice does not appear to undergo any deformation in spite of the minimal strain that must be overcome to match the TiO₂ epitaxy, indicating that the interaction between gold and titania is rather minimal. XPS measurements confirmed this lack of chemical interaction as Ti 2p and O1s did not show any core level shifts after deposition of gold for coverages above 3 ML (227).

On the anatase phase of titania the situation is somewhat different. HREM have revealed a strong epitaxial relationship between gold and the support due to the small misfit (0.9%)



Figure 13

Cross-sectional representation of $Au/TiO_2(110)$ interface viewed along (001]TiO_2 direction with (a) (111)_{Au}//(110)TiO_2 and (b) (112)_{Au}//(110)TiO_2 orientations. (258)

between the Au(111) face and the $TiO_2(112)$ plane (260). Although small gold particles have been theoretically predicted to favor icosahedral and decahedral shapes (206-210,261), the relationship between titania and gold is strong enough such that particles can grow in a Au(111) epitaxy up to rather large sizes (258). Giorgio and co-workers have shown with HREM that such a relationship also exists for small Au particles on anatase (110) (262). Comparing experimental data with simulated images, the best match was found when the particles were assumed to be strained such that their lattice constant was deformed to coincide with the support. The Au(110) was therefore assigned to grow along the TiO₂(110) face such that the Au-Au distances are contracted by 5.3 % along the Ti rows and expanded by 12.5 % between the Ti rows. Such a strong relationship was found to continue up to the fourth layer of gold.

The thermal stability of Au particles on TiO_2 has also been examined. Using HRSEM, Zhang *et al* observed substantial agglomeration of particles formed from a 3 ML dose of gold (5 nm average diameter) when annealing at 775 K (240). However, no encapsulation of gold by TiO_x under reducing or annealing conditions occurred, as has been observed for



Figure 14

STM images and particle size distribution of the Au/TiO₂(110) surface (a) before and (b) after CO:O₂ exposure (10 Torr, 2:1) showing Ostwald ripening. Images are 100 x 100 nm². (271)

platinum (263) and palladium (264) particles as a consequence of strong metal support interaction effects, so migration of gold into the oxide is not believed to occur. Previously Goodman's group had observed that gold particles deposited on a TiO₂ thin film at 120 K experienced substantial sintering between 120-400 K, but then reached a stable plateau that continued up to 800 K (265). Kitchin *et al* recently showed STM images that revealed significant sintering upon annealing of gold particles on TiO₂(110) to 773 K (266). The particle density was seen to drop dramatically and the average particle diameter increased from 0.45 nm to 0.75 nm. Kitchin *et al* also reported that the particles (after annealing) appeared to be strongly associated with defects on the TiO₂ surface.

Egdell and co-workers observed dramatic Ostwald ripening in their examination of the high temperature stability of small gold particles on $TiO_2(110)$ with STM (267). Simply said, Ostwald ripening is a growth/loss process by which large particles grow at the expense of small particles (268,269) usually resulting in a bimodal distribution. After deposition of 0.15 ML of gold at room temperature a fairly narrow particle size distribution centered around 27 Å was seen. After annealing the surface for seven hours at 750 K, the particle size distribution had a bimodal form with a center at 2 nm and a second center at 4 nm. The authors concluded that the Ostwald ripening must result from transport or migration of gold atoms from the edges of existing particles or which had not previously been incorporated into other clusters. This may be related to the low melting temperature of gold when in the form of small particles (270). Coalescence between particles at 750 K was also clearly observed in STM "movies". Although both of these growth processes may lead to catalyst deactivation it is difficult to assess which of these processes is dominant at the conditions of reaction.

Perhaps even more important than the initial nucleation and growth behavior of gold or its behavior under annealing conditions, is the behavior of supported gold under reaction conditions. Gold particles on titania were found to experience a form of Ostwald ripening when exposed to 10 torr of oxygen at room temperature (271).

Ripening in a reactive environment of O₂ and CO (10 torr, 2:1 CO/O_2) has proved to be even stronger. Clusters with rather narrow size distribution centered around a diameter of 2.6 nm and a height of 0.7 nm were observed to grow to 3.6 nm in diameter and 1.4 nm in height as shown in Figure 14. Furthermore, the presence of several smaller clusters of less than 1.5 nm in diameter was observed. Interestingly, when larger particles with an average diameter of 4.2 nm were exposed to 10 torr of oxygen at room temperature, essentially no ripening was observed. This indicates the ripening processes are connected intimately with the reactivity of the particles. The mechanism of the ripening process is currently unknown, although speculations surrounding the formation of a volatile oxide, as in the case of silver (272), seem unlikely. Willneff et al have recently presented evidence that when Au₂O₃ was exposed to CO at 400°C, an unidentified volatile species formed that is capable of transporting Au (273) although the oxide alone did not volatilize. When the surface of TiO₂(110) was pre-roughened with oxygen before gold deposition, the particles exhibited an enhanced stability in an oxygen atmosphere, possibly due to stronger binding of gold to surface irregularities. Gold does not seem to support the same type of titania regrowth that has been observed for palladium particles on $TiO_2(110)$ (274).

Of course the next step in observation of the sintering behavior of gold under reaction conditions is an in-situ measurement. Kolmakov and Goodman have examined gold on TiO₂(110) in a reactive environment of CO and O₂ with STM (275,276). An initially fairly uniformly sized group of particles was observed to undergo severe Ostwald ripening at 720 Pa and 450 K. As expected, increased surface temperature accelerated the ripening process. The authors suggested that the presence of oxygen served to weaken Au-Au bonds and thereby promoted sintering. Images taken under reaction conditions showed that the response of particles that were initially the same size could be quite different, as some particles decreased in size or even disappeared, while others seemed to remain stable. This result emphasizes the importance of the local environment in the structure and ultimately the reactivity of the particles.

AFM measurements by Iwasawa's group revealed significant sintering of Au particles on $TiO_2(110)$ under an

ambient atmosphere at temperatures as low as 363 K (277). However, pretreatment of the surface with ultraviolet radiation before deposition of gold resulted in a heightened resistance to agglomeration. The authors speculated that this behavior resulted not from surface defects but rather from the presence of hydroxyl groups which formed as a result of photo induced dissociation of water on the titania surface. The hydroxyl groups were suggested to react strongly with the goldphosphine precursor used by Iwasawa and thereby brought about the formation of highly stable small gold particles which showed limited agglomeration even at 493 K in air.

In an ex-situ experiment of gold on polycrystalline titania, Sykes *et al* observed extreme sintering of gold when exposing their sample to air or oxygen at room temperature for long periods (20 hours) (278). They concluded that the role of oxygen was to remove vacancies, which would otherwise bind the gold particles to the surface more strongly. Both Ostwald ripening and coalescence are thought to contribute to particle sintering.

Although most of the work on the growth, nucleation, and thermal stability of gold has been performed on the $TiO_2(110)$ surface, gold has been studied on a variety of other oxides. Early TEM work by Poppa and others demonstrated that gold particles on mica (279-281), MgO (282-287) and sapphire (288) were quite mobile and that vapor deposited particles would move rapidly on the surface in an attempt to find a favorable epitaxial orientation such as (111)_{Au}//(001)_{Ma0} or $(110)_{Au}//(110)_{MqO}$. Green *et al* found that particles grew epitaxially on vacuum cleaved MgO crystals, but that on air cleaved crystals the orientations were random until thick (100 Å) films were grown (283). Increases in the particle density were seen when the MgO substrates were sputtered with Ar ions prior to gold deposition as expected (286). However, attempts to induce similar nucleation sites with an electron beam only resulted in an increased nucleation density on MgO(111) and not on MgO(100), where an unexplained reduction in the nucleation density was observed.

More recently Henry's group has re-examined growth of gold particles on mica (289) and MgO (290,291) with AFM. Large particles grown on mica exhibit the Au(111) face as shown in Figure 15 with a very flat topography (289). The 23 $x\sqrt{3}$ reconstruction has not been observed on the top facets, although this is probably because the long-range periodicity of the superstructure is large compared (6.5 nm) to the size of the particle (22 nm average). The sides of such flat hexagonal or triangular particles displayed a lattice constant characteristic of the Au(100) surface. Gold particles deposited on MgO microcubes were found to grow in a $(001)_{Au}//(001)_{MgO}$ epitaxial relationship and exhibited fcc structures up to 4 nm with a expanded lattice parameter of 2.9% (290). Particles of greater than 4 nm diameter were seen to adopt a $(111)_{Au}//(100)_{MgO}$ epitaxy as well, but the

lattice constant observed in the larger particles was expanded to that of bulk gold regardless of orientation. Ajayan and Marks had previously observed that the structure and orientation of very small gold particles (d ~1 nm) can fluctuate significantly even at room temperature as the energy differences between conformations is on the order of kT as has been seen in calculations for gas phase clusters (292). However, the authors explain that the substrate can play a critical role in the stabilization of the particle in particular states. Giorgio et al ascribe their lack of observation of such transitions between structure, known as "quasimelting", to the high interfacial energy which must be overcome when the particles are epitaxially oriented. On MqO(100), Højrup-Hansen et al examined the nucleation behavior of gold and found that as the deposition time increased, the cluster density rapidly reached a saturation plateau (291). For long deposition times, the cluster density began to decrease once again due to coalescence. The saturation density was found to decrease with increasing temperature and by means of an Arrhenius relationship determine an activation energy for nucleation to be 0.12 eV. The authors concluded that since their results could not be described by a homogeneous nucleation mechanism, nucleation on point defects must be the prevailing process.

Robins and Rhodin found that the nucleation rate was directly proportional to the deposition rate leading them to conclude that nucleation was dominated by point defects (293). Furthermore, their observation that the nucleation density reached a constant maximum regardless of deposition rate or substrate temperature was presented as additional proof that the nucleation process was defect mediated. However, this result is in contrast with Henry's finding of an Arrhenius type relationship between nucleation density and substrate temperature. In Robins and Rhodin's model, point defects acted as perfect sinks for nucleation but recent theoretical (294) and experimental (295) work has shown that this does not apply for high substrate temperatures. The authors were able to observe a number of different types of defects on the MgO surface, all of which acted as nucleation centers for gold (296).

In their study of a gold particle grown on a MgO(100) step, Kizuka and Tanaka saw that the particle could rearrange continuously to maintain the lowest energy structure when holding the sample substrate at room temperature (297). Examining the ratio of Au(100) facets to Au(111) facets, repetitions in the truncation pattern were seen to emerge as the cluster grew until reaching a size of over 1500 atoms at which point rigid epitaxy with the support is lost. Pauwels and co-workers found in their examination of HREM images that both the MgO surface layers and the Au(100) layers were strained around the interface in an effort to match lattice constants (298). Interestingly, the lattice spacing at the edges of the particle were expanded to compensate for the compression of the lattice constant at the particle interior. In their examination of Au on MgO using TEM, Blick and co-workers found that gold grew two-dimensionally at low coverages forming thin rafts before exhibiting more three dimensional growth at higher coverages of gold (299).

Growth of gold on silica thin films has been studied by Luo *et al* (300). Similar growth behavior to Au/TiO₂(110) was observed as the growth mode switched from 2-D to 3-D at approximately 0.1 ML. For gold deposited at 100 K, sintering was observed to be strong when annealing the sample to 600 K. However, above 600 K sintering slowed until gold was finally desorbed from the surface at above 1000 K. When comparing LEIS data for the Au/SiO₂/Mo(110) model catalyst with the Au/TiO₂/Mo(100) model catalyst, in both cases the Au signal lost about half its intensity between 100 K and 800 K, indicating diffusion behavior of gold is essentially the same for both surfaces (265,300).

However, unlike on $TiO_2(110)$, growth of gold on thin alumina films has been seen by ourselves and others to be 3-D even at very low coverages (301,302). Well ordered alumina films can be grown through the oxidation of the surface of a NiAl(110) single crystal. The thin alumina film, which we have studied extensively, exhibits large terraces, on which line and point defects exist, which may act as nucleation centres during metal deposition. STM images of gold deposited at 300 K showed that gold particles are randomly distributed on the surface. Based on previous extensive studies on metal nucleation and growth on alumina (233-235), the absence of preferential nucleation for deposition at 300 K indicates that the gold atoms have a fairly low mobility on the alumina surface. Even at a coverage of 0.05 ML of gold, particles were seen to exhibit a hemispherical particle shape, indicating 3-D growth from the onset. The Au particles formed on alumina showed a narrow size distribution. In addition, the system morphology was quite stable towards heating up to 600 K: the particles only slightly sintered as evidenced by STM and LEED measurements.

Carrey *et al* have proposed that large gold particles on an amorphous alumina film can be more mobile than single atoms or small clusters (303,304). The authors observed that the nucleation density of gold was not affected strongly by deposition temperature leading them to conclude that the nucleation must be dominated by defects. Furthermore, the deposition resulted in a bimodal size distribution, indicating that particles above a certain size could more easily diffuse and coalesce during deposition while smaller particles were trapped on defects.

The thin FeO(111) (305,306) film grown on a Pt(111) single crystal has only recently begun to attract attention as a suitable oxide substrate for model metal/oxide systems



Figure 15

(a) AFM topographs (b) lateral force and (c) normal force images of the top face of a flat gold particle formed on mica. (Image size 6×6 nm²). (289)



Figure 16 High resolution STM image of a monolayer gold particle formed on a thin FeO(111). (Image size $10 \times 6 \text{ nm}^2$).

(302,307). The oxide layer exhibits an O-Fe-Pt-Pt- stacking and strongly relaxed surface, with O-Fe and Fe-Pt distances being \sim 0.6 and \sim 1.8 Å, respectively. The surface shows a characteristic superstructure, clearly visible in STM and LEED, formed by a \sim 10% mismatch between FeO(111) and Pt(111) lattices (305).

Gold deposition first resulted in decoration of steps and "holes" present in oxide film, which typically cover less than 3% of a monolayer. Subsequently, small Au clusters started to nucleate and grow at the terraces as well. At high coverages, Au aggregates were essentially homogeneously distributed on the FeO(111) surface. The nucleation density of gold on the FeO film was approximately half that of gold on Al₂O₃/NiAl(110). This implies that, albeit both oxides are very thin (< 5 Å) and terminated by oxygen, gold diffusivity on these supports is quite different. The higher diffusivity observed for gold on FeO probably results from a lower defect density on the FeO film relative to alumina.

The high resolution STM image presented in Fig. 16 shows a 1.5 nm diameter island of just a single (111) layer which is preferentially formed for ~ 0.1 Å-thick Au overlayer. As both

the support and Au island are atomically resolved, it is clear that the gold layer is in good epitaxial relationship with the oxide support. With increasing amounts of gold, threedimensional particles up to 7 nm in diameter and 1.5 nm in height could be grown (for 2 Å-thick overlayer). Since good epitaxy with support was observed for Au monolayer islands, it seems likely that large Au particles grow such that they expose (111) top facets, as has been observed for gold particles on mica (289).

We have extended the studies to other iron oxide films. Previous studies of the $Fe_3O_4(111)$ films formed on Pt(111) have shown that the regular surface is terminated by 1/4 of a monolayer (with respect to oxygen sub-layer) of Fe^{3+} cations (306). Besides terrace steps, the defect structure, which may influence the nucleation and growth of Au particles, includes iron vacancies and, in some cases, small domains exhibiting "biphase ordering", whose presence at low concentration can be only observed with STM (308,309).

Accordingly, the gold diffusion on $Fe_3O_4(111)$ was found to be limited relative to FeO, thus preventing sintering at 300 K. However, heating to higher temperatures promoted sintering such that particles annealed to 500 K grew and displayed better faceting (302). The histogram analysis of STM images for annealed samples revealed that height of the particles was found to be by multiples of 2.5 Å (equal to the height of a monolayer of gold in the Au(111) orientation) which implies that particles grow by increasing the number of the layers in the cluster. In addition, the annealed particles are well ordered exhibiting mostly hexagonal (and trigonal) shape. It seems plausible that the top facets of the particles show the (111) surface owing to small (~ 3%) lattice mismatch between Au(111) and Fe₃O₄(111) surfaces.

The STM results for the α -Fe₂O₃(0001) film are very similar to those found on Fe₃O₄ films. Once again, 3-D Au particles are found homogenously distributed on the oxide surface, and they exhibit a hemi-spherical shape. Previously, Haruta found by TEM that gold forms hemispherical particles on Fe₂O₃ with an epitaxial orientation of (111)Au//(110)Fe₂O₃ (310). However, the Fe₂O₃ surface termination is critical to the preparation conditions (306), and therefore the nucleation behavior is currently a subject of investigation.

5.2 Electronic properties of supported Au particles

It is well known that small particles supported on oxides can exhibit electronic, optical, and magnetic characteristics different from bulk materials (311-316). In the case of gold, this is especially important as several researchers believe it is precisely these changes in the electronic character that give rise to its unique catalytic properties. Although gold catalysts are not thought to exhibit Strong Metal Support Interactions (317), at least in the classical sense (318), perhaps the role of the



Figure 17

Turnover frequency and band-gap measured by STM/STS as a function of the diameter of Au islands formed on TiO₂. (2 (data from 192))

support is to influence the electronic nature of the gold through charge transfer. Aside from specific support effects, quantum size effects (319,320) are also frequently invoked to explain the exceptional activity of gold nanoparticles. Valden et al demonstrated with Scanning Tunneling Spectroscopy that the maximum in activity for Au particles on TiO₂(110) coincided with the onset of an appearance of a band gap, marking a transition point from bulk metal to a semi-conductor (192) as shown in Figure 17. At a diameter of 3.2 nm and height of 1.0 nm (~ 300 atoms) a band gap of 0.2-0.6 eV can be measured which increases up to 1.4 eV were measured for the smallest particles (d < 2.0 nm). Valden et al asserted that the gold particles experience this unusual activity as a consequence of their reduced dimensionality and suggested that particle thickness may be the key parameter. Similar measurements of band gaps for Au particles on HOPG indicated that the metalnon-metal transition took place at even smaller particle sizes (~ 1 nm) (321). However, this could be explained by differences in geometry between particles on various supports as small gold clusters on HOPG were found to be spherical (322).

In their study of acrolein hydrogenation over Au/TiO₂ and Au/ZrO₂ catalysts, Claus and co-workers also invoked quantum size effects to explain a change from the preferential hydrogenation of C=O to C=C as the particle size decreased below 2 nm (323). Small gold particles sitting above oxygen vacancies of the oxide may be electron rich due to charge transfer from the support and would therefore be predicted to experience a repulsive interaction with the C=C bond. However, due to the quantum size effects, the *d*-band of a small gold particle will possess a decreased density of states which lowers the repulsive interaction. At the same time, the depletion of the *d*-band also results in a loss of back donation into the C=O bond.

Therefore since the activity of gold has been linked to the transition from metallic to non-metallic particles, the exact point at which this transition occurs is the subject of much interest. Using size selected clusters supported on amorphous carbon, Berry and co-workers found that Au₃₃ (and smaller clusters Au₂₇, Au₇, and Au₅) showed significant shifts in both the valence and core regions by XPS (324). The authors correlated these shifts with the average coordination number of the atoms in the cluster. From an extrapolation of their data, the transition from metallic to non-metallic behavior was predicted to occur at around 150 atoms. In a similar examination based upon the 5d band splitting, Lee etal had estimated that this transition should take place at a particle diameter of more than 1.9 nm, which corresponds to a hemispherical particle with over 100 atoms (325). Using HREELS, Guo et al. estimated that this transition occurred for gold on TiO₂ (110) at a coverage of 0.2 ML Au, which corresponds to particles of about 2 nm in diameter and 2 atomic layers high (242). For gold on a thin alumina film, the transition was not seen until a coverage of 0.3 ML of Au (242). Whetten and co-workers found that discrete states in the optical absorption spectra began to emerge at diameters below 2 nm (~ 200 atoms) (326). The authors noted that this transition took place at the same point where a structural transition occurs, suggesting the two phenomena may be related (207). Stievano et al employed Mössbauer spectroscopy to observe changes in the electronic character of small gold particles supported on Mylar as the dimension was shrunk (327). Measurements revealed an asymmetry as the particle size decreased below a 1.5 nm diameter, due to the increased fraction of surface atoms. A decrease in the ion splitting was explained by an electron transfer from the core to the surface atoms. Recently, Boyen et al showed by XPS that Au particles on diamond displayed the valence band spectra as bulk gold down to a diameter of 1.6 nm (~ 100 atoms) (328). Lin et al used the field emission energy distribution to observe the metal-non-metal transition for gold (329). Particles below 3.0 nm (approximately 800 atoms) were observed to display discrete electron energy states as opposed to the continuum of states in bulk gold. Calculations based on an electronic shell model were in good agreement with the experimental observations.

Such examinations underline the importance of photoelectron spectroscopy in studying the electronic character of small gold particles. Core level binding energy shifts can be interpreted as initial state effects which result from quantum size effects inherent to the particles themselves and final state effects which result from relaxation processes and are strongly connected to the support (234, 330). Of course, smaller particles also have more surface atoms, which due to their decreased coordination results in a shift of the *d*-band toward the Fermi

level and a corresponding decrease in the binding energy of the core electrons. In the case of gold, Wertheim and coworkers have measured this surface state shift to be ~0.4 eV (331). For small unsupported gold clusters Häberlen *et al.* calculated that shifts to lower binding energies can range up to 1.0 eV for Au₁₃ clusters and that even Au₁₄₇ clusters show an overall shift of 0.4 eV (206). Furthermore it is possible to distinguish between initial and final state effects through analysis of the so-called Auger parameter (332) by which one can determine to what extent the relaxation effects play a role in the binding energy shift by the creation of a double hole. Photoemission experiments are also useful in observation of charge transfer from the support to the particle (or vice-versa).

Mason attributed positive binding energy shifts observed for Au/SiO₂ and Au/C to an initial state effect arising from changes in the electronic configuration of small particles due to rehybridization of the valence band (333). Mason asserted the relative similarity in the binding energy shift for gold on two very different substrates was strong evidence for the importance of initial state effects. In a similar comparison, Liang *et al* examined the valence spectra of gold deposited on aluminum and aluminum oxide and saw similar behavior with respect to the position of the *d*-band and the degree of *d*band splitting on both substrates (334). This led the authors to conclude that the deviations from bulk behavior must arise from intrinsic electronic properties of the small particles.

In contrast, Wertheim *et al* concluded that such positive binding energy shifts must arise from final state effects from their comparison of XPS spectra of Au particles on vitreous carbon (non-conducting) and Metglas (conducting) (335). A slight negative core level binding energy shift was observed in the latter case, even though the characteristic narrowing of the *d*-band was observed in the valence spectra indicating the formation of small particles. Kim and Winograd used implantation of Ar into gold and gold supported on SiO₂, as a way to compare the chemical environment of gold in the bulk with gold on SiO₂ (336). The positive binding energy shift of Au 4*f* for small gold particles on silica was interpreted as a final state effect as Ar was found to undergo a shift of similar magnitude when compared to Ar implanted in a bulk gold sample.

Although Coulthard *et al* acknowledged that their observation of positive binding energy shifts was consistent with the idea of hybridization effects endorsed by Mason (333), XANES measurements indicated that small gold clusters possess an enrichment of d charge (337). As an initial state view would predict a *d* count in between 9 (single atom) and 10 (bulk metal), this appears inconsistent with the XANES data. Since XANES experiments do not suffer from charging effects, one can conclude that the shifts observed by XPS must arise from final state effects.

Of course, both initial state effects and final state effects will be present and several researchers have found that one must consider both effects to obtain an accurate picture for the origin of binding energy shifts. When Cordes and Harsdorff examined the 4*f* region of gold particles supported on amorphous carbon, they found that the positive core level shift had disappeared for particles with a diameter greater than 1.5 nm (338). However, the valence band region did not exhibit bulk behavior until the particles were at least 3.5 nm in diameter. Thereby, the authors concluded that the same mechanism cannot be controlling the deviation from bulk behavior in the two separate energy regions. Cordes and Harsdorff hypothesized that while the core region spectra are dominated by relaxation processes, the valence band depletion was a result of decreased spin-orbit coupling in smaller particles. Roulet et al examined the valence region of gold particles from 1.5-6.0 nm on NaCl and also observed depletion near the Fermi Level as the particle size was decreased (339). Particles of 4.0 nm or less exhibited significant band narrowing as well. The band narrowing was interpreted as a consequence of reduced coordination.

In a comparison of particles with different morphologies, but approximately the same total number of atoms, Costanzo *et al* observed that two-dimensional gold particles grown on HOPG experienced smaller binding energy shifts as compared to more spherical particles deposited on amorphous graphite (340). While the positive binding energy shift was attributed to final state effects, the differences between particles were interpreted as initial state effects arising from the higher percentage of surface atoms in the particles on HOPG.

In Dalacu *et al's* examination of gold particles of various sizes on SiO₂ supports with XPS, positive binding energy shifts for the Au 4*f* peak of up to 0.5 eV were observed for the smallest particles (341). After annealing, the shift increased to even higher binding energies (up to 1.3 eV shift), which was interpreted to be primarily a final state effect resulting from decreased contact with the support with only a minor contribution by initial state effects due to a decrease in the number of exposed surface atoms.

In an examination of gold on ZrO₂, Zafeiratos and Kennou used the Auger parameter to separate initial and final state effects (342). While final state effects dominated the overall binding energy shifts observed, it was possible to observe initial state effects reflecting the average coordination number up to a gold coverage of 1.5 ML. It should be remembered, however, in analysis of the Auger parameter that the relaxation comparison of the double hole and single hole should be from the same shell. Recently, in our own analysis involving copper and cobalt particles (343), we have obtained evidence that lattice contraction in very small particles can induce positive binding shifts due to hybridization as predicted by Mason (333). This situation is of particular importance to gold which has a strong tendency to undergo contraction in the surface layer (206,344-348), hence the reconstructions of Au(111) and Au(100).

In an investigation of support effects, Chusuei *et al* measured binding energy shifts of gold on SiO₂ and TiO₂ and saw rather large differences between the two substrates (0.8 eV for Au/TiO₂, 1.6 eV for Au/SiO₂) (349) as shown in Fig 18. Due to a lack of structural data for Au/SiO₂, it is not possible to say precisely how these effects originate. However, this data seems to indicate that interactions of the support with gold particles are present that must affect the electronic character of the particles. Band bending observed by shifts in the XPS for Ti 2p (0.15 eV) for submonolayer coverages of gold on titania, give support to the notion that the positive binding energy shifts observed in the Au 4f result from the interaction between gold and the oxide support (227).

Yang *et al* have calculated that the TiO₂ support should induce negative binding energy shifts of 1.1-1.5 eV for a gold monolayer lying above it due to transfer of charge from the support to the particle (248). At first this result appears inconsistent with Chusuei *et al* (349). However, as the authors point out, Au particles on titania do have a negative core level shift as compared to Au/SiO₂ (which is a non-interacting support). Some researchers have invoked final state effects as an explanation for the discrepancy between theory and



Figure 18

Plots of binding energy measured by XPS of the Au 4f7/2 core level as a function of Au cluster coverage on (a) $TiO_2(110)$ and (b) SiO_2 surfaces. (349)

experiment. However, in recent DFT calculations it has been demonstrated that final state effects are insufficient to explain the positive binding energy shifts observed by Goodman's group and others. Yang and Wu explain the positive binding energy shifts observed through the interaction of gold with oxygen vacancies (350).

Wu and co-workers went on to perform calculations for adsorption of gold on MgO(001), taking a special interest in perturbation of the electronic structure by defects (254). Like on TiO₂(110), gold on MgO(001) was found to experience core level shifts to lower binding energies for adsorption on the regular surface. On defects (oxygen vacancies), the situation was reversed as gold was found to undergo a shift of 1.8 eV to higher binding energies. These calculations seem at odds with XPS experiments for Au/SiO_2 and Au/TiO_2 showing that reduced surfaces give rise to negative shifts in the binding energy as one would expect for an anionic gold species (351,352). Howard et al examined the influence of defects on the support upon the electronic character of gold particles (352). After bombardment of the TiO₂(110) surface with argon before gold deposition, only very small core level shifts were observed (0.1 eV at 0.5 ML Au). Therefore the negative shift due to charge transfer from the Ti³⁺ states to gold balanced out the shift to higher binding energies due to final state effects that was observed for particles on the defect free surface. Even more dramatic, after deposition of gold on the reduced surface, the signal for Ti^{3+} was found to practically vanish. Examination of the valance band uncovered a strong depletion around the Fermi level for small particles due to either initial state effects or charge transfer to the substrate. On the defective surface an even greater shift in the onset of photoemission was observed, leading the authors to conclude that the depletion of intensity near the Fermi level resulted from an interaction between Ti 3d defect states and gold. Vijay et al in their calculations of the growth of gold on titania found that the charge left behind by a vacancy was not isolated on the two exposed titanium atoms but rather was spread over a fairly large ensemble in the oxide (250). Therefore, one should consider that vacancies, which previously had been thought to be a highly localized phenomenon, may have long-range effects on the surface chemistry.

Goodman and co-workers found in their calculations a shift in the valence band for Au/TiO₂ towards the Fermi Level by 1.8 eV for a single monolayer of gold (248). This shift in the 5*d* band toward the Fermi level suggested that small supported gold clusters may not be so "noble" and should therefore have enhanced chemical adsorption properties as compared to bulk gold. Lopez and Nørskov found evidence in their calculations that this result was dependent upon the coverage of gold (253). For coverages of less than one monolayer, the *d*-band was shifted to the Fermi level by



Figure 19

Photon-emission spectra of Au particles of various sizes on $TiO_2(110)$ as marked in the STM image shown in the inset (size 130x130 nm²). The emission is dominated by decays of Mie-like plasmons. (353)

upwards of 1.0 eV when compared to bulk gold. However, such initial state effects were balanced by a strong interaction between gold and the underlying support, leading to a shift in the position of the *d*-band to higher binding energies with respect to a free gold cluster. As the coverage of gold was increased, the fraction of highly coordinated gold atoms in the particle increased, leading to more metal-metal bonding and thereby drawing electron density away from the particle-support interface. For coverages as low as two monolayers, the density of states was essentially identical to bulk gold.

Wang and Hwang also observed a shift upward in the DOS when Au was placed upon a titania surface, indicating there should be considerable changes in the chemistry of supported gold compared to bulk gold surfaces (251). Perhaps surprisingly, the strong interaction between gold and a defective TiO_2 surface, actually serves to broaden the 5*d* states of gold, possibly reducing the particle's reactivity. However, charge density plots also reveal some charge transfer from the defect site to Au.

In our own lab, we have examined the electronic properties of small gold particles using photoemission STM (353,354). The Mie plasmon resonance energy of a cluster can be probed by injecting an electron into the cluster with an STM tip and analyzing the photons emitted. In a simple synopsis, the Mie plasmon is the collective oscillation of the conduction electrons in a particle relative to the core. The photoemission spectra of 7 nm diameter Au particles supported on a $TiO_2(110)$ single crystal revealed that the electronic character of the particle was essentially unaffected by the support. The resonant photon energies were found to increase with decreasing particle size as the contribution of surface atoms with more localized *d*-electrons increases as

shown in Figure 19, although particles in the non-metallic range could not be probed due to interference from tip induced plasmons. This is a consequence of the peculiar dielectric function of gold, which passes through a plateau region precisely in the range of the Mie plasmon resonance for gold and results in a heightened sensitivity of the emission to the local environment (i.e. the STM tip condition).

5.3 Adsorption and reaction on supported Au nanoparticles

In some of the earliest work, Haruta et al (9,191,355) identified that the onset of activity of gold catalysts for CO oxidation could vary widely with the choice of support. Even in a comparison of different forms of titania (anatase, rutile, amorphous), large differences in reactivity can be seen (356). Some researchers ascribed this effect to differences in the particle size distribution on the various supports (although the gold loading was kept constant) (357,358). However, Grunwaldt et al showed a large difference in activity between zirconia supported gold catalysts and titania supported catalysts (Au/TiO₂ >> Au/ZrO₂) even though the particle sizes were identical (359). The authors believe that the support affected the gold particles' ability to activate (dissociate) oxygen. The results were interpreted as differences between reducible and non-reducible supports. Behm and co-workers (226) found that the activity of gold catalysts supported on "non-reducible" metal oxide supports was considerably less than that of "reducible" metal oxide supports, as Au/Fe₂O₃ and Au/TiO₂ were over five times more active than Au/Al₂O₃. Furthermore, Behm's group reported that the activity of these inert supports was more sensitive to the size of the supported particles. The authors ascribed this difference in activity to a possible difference in the mechanism of oxygen adsorption. More recently, the oxidation of CO by gold supported on alumina and titania has been compared by Haruta and coworkers (360) and only slight differences in activity between gold catalysts were found when care was taken to ensure that the gold particles are well dispersed.

Supported gold particles have shown in numerous examples to exhibit heightened interaction with adsorbates as compared to single crystal surfaces. For example, using XPS, Rodriquez *et al* observed the dissociation of SO₂ on 2.6 nm diameter gold particles on TiO₂ (110), even though SO₂ adsorbs only very weakly on the bulk surfaces of gold (252). While SO₄ was found to be the only product on the clean TiO₂(110) surface, SO₂ experienced total dissociation when in contact with gold on titania. XPS experiments also showed an increase in the relative fraction of Ti^{(4-x)+} after annealing to 750 K when gold was present as compared to its absence. The authors hypothesized that gold served to increase the rate of vacancy exchange between the surface and the bulk of TiO₂, while at the same time gold is electronically modified

by the presence of oxygen vacancies. Collaborative density functional theory calculations revealed that SO₂ dissociation was greatly enhanced on gold when the particle is located above an oxygen vacancy.

Rodriguez and co-workers found further evidence of support effects in their comparison of SO_2 adsorption on Au/TiO₂ as compared to Au/MgO (361). In the case of MgO as a support, no dissociation of SO_2 was observed, although the desorption temperature was much higher than that of SO_2 from an Au single crystal. This would imply that small gold particles on MgO are a somewhat intermediate case: less active than Au/TiO₂ but more active than bulk gold surfaces. Since the Au particle size was expected to be about the same on both surfaces, Rodriguez *et al* attribute the differences observed to metal-support interaction effects.

As discussed in the previous section, the electronic character of gold may be perturbed by an interaction with the support. A consequence of the perturbation may be unique adsorption behavior that neither gold nor the support material exhibit alone. An example of this type of synergism is the dissociation of H_2S on Au/Cr_2O_3 catalysts as studied by Rodriguez and co-workers with synchrotron radiation measurements (362). *Ab initio* calculations supported a mechanism whereby a charge transfer occurred from the gold particles to the oxide support and H_2S decomposed easily on the electron deficient gold.

Even when gold is deposited on a "non-interacting" support such as activated carbon, support effects can be seen. Bianchi *et al* observed differences in activity of various Au/C catalysts for the oxidation of ethylene glycol depending upon the support although the particle sizes were essentially equal (363). The differences in activity were ascribed to presence of phenolic groups on the surface and their interaction with dispersed gold.

The synergism between gold and the support can also be observed by changes in the metal oxide. For example, it has been seen that certain metal oxides like CeO₂ (364) and Fe₂O₃ (365) are more easily reducible in the presence of gold. Using ESR spectroscopy, Andreeva *et al* found that the V=O bond length was lengthened in Au/V₂O₅ catalysts after gold deposition leading to a higher delocalization of 3*d* electrons and suggesting a strong interaction between Au and the support (366).

Given the complicated nature of the relationship between gold and metal oxide supports, it seems logical to begin our examination of size and support effects in gold catalysis with a discussion of the simple case of CO adsorption on gold model catalysts like those described in section 5.1. In Rainer *et al's* examination of the adsorption of CO on an Au/Al₂O₃ thin film model catalyst with IRAS, CO was found to exhibit a single peak at 2100 cm⁻¹ at 100 K for 0.5 ML Au deposited at 300 K (367). In agreement with Ruggerio and Hollins work on Au(332) (83) this peak underwent a reduction in intensity and a blueshift of 11 cm⁻¹ as the sample was heated to 200 K. When higher gold coverages were used, a small blue shift to 2110 cm⁻¹ was observed. The former effect is in contrast to the behavior of CO on other metal surfaces such as Pt (93) and Pd (94) and is ascribed to the unique electronic character of gold by which the back donation of electrons from gold into the 2 π^* orbital of CO is reduced (368). The latter effect is understood as a consequence of a decrease in the density of adsorbed CO as the particle size is increased. Winkler et al found similar results in their examination of CO on Au/Al₂O₃/NiAl(110) using IRAS (369). In accompanying TPD experiments, CO was seen to desorb in two peaks at 105 K and at 145 K from gold particles of ~10 atoms. Attempts to observe oxygen desorption or CO₂ production on these small particles failed.

In our work examining CO adsorbed on Au particles on an Al₂O₃ thin film, TPD experiments revealed two clearly resolved peaks, with the first peak at 120 K and another located at a higher temperature, depending on the nominal thickness of the Au overlayer (302). Bearing in mind that CO does not adsorb on the alumina film above 77 K, observation of two desorption states may, in principle, indicate separate CO adsorptions on the metal and at the metal-oxide interface. However, based upon Ruggerio and Hollins observation of CO desorption from the stepped Au(332) single crystal surface (83) we conclude that both peaks could be attributed to desorption from gold particles. The position of the high temperature peak gradually shifts from 210 K to 160 K as the Au nominal thickness increases from 0.1 to 1.3 Å. This result clearly shows a particle size effect such that small Au particles adsorb CO more strongly.

The intensity of CO desorption signal reached saturation at a fairly low coverage of gold, as the spectra are nearly identical for the samples possessing 1 and 5 Å-thick Au overlayers. This finding can be understood if CO only adsorbed on low-coordinated Au atoms, but not on the regular terraces (see results for Au/FeO below), in good agreement with theoretical predictions (190). Therefore, it seems likely that gold deposition beyond 2 Å thickness leads only to the gradual development of the flat terraces on the gold particles, which are inert towards CO.

When examining the TPD data of CO adsorption on the Au/FeO samples as deposited at 77 K, a strong particle size effect was seen, with desorption states observed at temperatures of up to almost 300 K for the lowest gold coverages (smallest particles) (302). After annealing to 500 K, a reduction in the signal was observed and the size effect was completely suppressed, as TPD spectra showed two peaks at 130 and 200 K, independent of the amount of gold deposited. This finding indicates that the particles after annealing have reached some minimum critical size beyond



Figure 20

IRAS spectra of CO exposed to Au/FeO surface at 90 K as a function of Au coverage. (a) as deposited particles at 90 K (b) after annealing to 500 K

which their adsorption behavior is size independent. However, particle size is not the only variable to consider as restructuring has likely taken place simultaneous to the sintering process and the low coordinated gold atoms which have been theoretically predicted to adsorb CO most strongly may have disappeared due to restructuring. In fact, we find that even monolayer islands formed after annealing to 500 K, as seen in Fig.16, display the same CO adsorption behavior as large particles or gold single crystals (370). As the gold particles are known to nucleate on regular lattice sites as opposed to defects (302), we can exclude support effects. Therefore, we can conclude that size effects are only present in our system as such that smaller particles, particularly before annealing, contain a higher percentage of highly uncoordinated gold atoms which favor CO adsorption. In turn, we assert that quantum size effects related to particle thickness as proposed by Valden et al (192), are not likely to be a significant factor in CO adsorption.

Figure 20a shows IRAS spectra taken at saturated CO exposure for samples deposited at 90 K as a function of gold nominal thickness. At a coverage of 0.1 Å, a feature at 2166 cm⁻¹ and a second at 2131 cm⁻¹ are observed with approximately equal intensity. With increasing coverage, the higher frequency peak (2166 cm⁻¹) decreases before disappearing at 2 Å Au, while the second feature grows and shifts rapidly to lower wavenumbers from 2131 to 2108 cm⁻¹ (371). This trend is different than that observed by previous researchers examining gold particles on thin alumina films,

where CO adsorbed on the smallest particles was observed to be at lower numbers (~2100 cm⁻¹) and blue shifted to 2110 cm⁻¹ as the particle size increased (366,368). The origin of the difference between our system and these examples is unclear, but Goodman's group has recently studied CO adsorption on Au particles on a TiO₂ thin film and also observed adsorption frequencies higher than that of CO adsorbed on bulk gold, which they attributed to a support effect of unknown origin (372).

As shown in Fig. 20b, annealing to 500 K results the observation of a single state at 2108 cm⁻¹ independent of the Au coverage with the disappearance of the state at 2165 cm⁻¹, just as in the case of high gold coverage. Bearing in mind the sintering effects of annealing, the peak at 2108 cm⁻¹ can be straightforwardly assigned to CO adsorbing on metallic gold. We propose that the species exhibiting the peak at ~ 2165 cm⁻¹ is probably CO adsorbed on very small clusters which can undergo charge transfer due to a specific interaction with the support initially on defects (371). Since at low coverages, the nucleation and growth is strongly influenced by steps and holes present on the FeO film (302), it seems likely that such charged species could be present. In previous IRAS studies on different catalysts systems like gold supported on zeolites (373), Au/ZrO₂ (374), Au/TiO₂ (374-376) and Au/Fe₂O₃ (377,378) similar CO stretching bands frequencies have been also detected and assigned to CO adsorbed on Au⁺ species or Au interacting with hydroxyl groups on the surface (see Table 2). However, in our experiment, this feature appears after the main peak observed at ~ 2108 cm⁻¹ and therefore implies that CO adsorbs on such species more weakly relative to the metal clusters as has been seen by Fan et al (379), although this is in apparent contradiction with the view of some investigators that charged gold adsorbs CO more strongly (373,378,380-382). Interestingly, the single peak observed in the annealed samples did not shift and simply grew monotonically at increasing Au coverage, supporting our finding from TPD experiments indicating no apparent change in the possible adsorption states for CO on monolayer islands as compared to large particles. Therefore, we propose that CO adsorption on gold particles only includes the low coordinated surface atoms. This result can be traced back in order to explain saturation behavior for CO adsorption on gold deposited on alumina at the high coverages.

Gold supported by Fe_3O_4 is of particular interest due its proposed role in the water-gas shift cycle of Au/Fe₂O₃ catalysts (11,383). It has been proposed that the dissociative adsorption of water on gold leads to a spillover of active hydroxyl groups onto the ferric oxide. The iron oxide surface is then reduced to Fe_3O_4 by the formation and decomposition of formate or carbonate intermediate species before subsequent reoxidation by dissociative adsorption of water. Furthermore, it has been shown by XPS that the surface of Au/Fe₂O₃ catalysts can actually be dominated by Fe_3O_4 (384).

Figure 21

Comparison of CO TPD spectra for Au particles deposited on different oxide films showing that CO adsorption is essentially independent of the support at a given particle size (~ 3 nm)

The CO adsorption studies of Au/Fe₃O₄ were more complicated relative to FeO and alumina films by the fact that the magnetite surface may adsorb CO and possesses desorption states which partly overlap with those of gold. After subtraction of the signals from the clean support, the CO TPD spectra showed the size effect as seen for other supports, i.e., the smaller the particles, the stronger the CO-Au interaction (302). In contrast to the FeO(111) support, the effect was partly retained for Au/Fe₃O₄(111) even after heating to 500 K. However, strong reduction of the CO desorption intensity clearly indicated the morphology changes upon annealing. For CO adsorbed on Au/Fe₂O₃, the behavior is quite similar. The CO TPD spectra for "as deposited" at 90 K samples show a desorption state extended to ~300 K, which disappear after annealing to 500 K.

The CO adsorption studies on Au deposited on alumina and iron oxide films performed in our group clearly showed the particle size effect in that small particles adsorb CO more strongly. For the smallest Au coverage, CO may desorb at temperatures close to 300 K, which is in the temperature range of the working Au catalysts for low temperature CO oxidation and has never been observed for Au single crystals. As discussed above, the presence of such a state is likely due to the presence of highly uncoordinated gold atoms.

In Fig. 21 CO TPD spectra obtained for gold deposited on various supports are compared for particles that have been annealed to 500 K. The comparison clearly points out that, for a given particle size (ca. 3 nm as measured by STM), the interaction of CO with gold particles on different supports is essentially identical. Therefore, the results show that support effects frequently observed for CO oxidation on real catalytic

Table 3

Results of CO adsorption on various gold surfaces.

Surface	Particle size	Condition (CO pressure)	Frequency observed	Assignment	Ref.
Au/SiO ₂	5 nm	0.3-84 torr @ 300 K	2120-2110 2145	Au ⁰ oxygen covered Au	542
Gold film		1 ML @ 113 K	2110		81
Au/MgO	8 nm	1-15 Torr @ 300 K	2060-2040	multi-bonded CO	444
Au/SiO2	40 nm	30 Torr @ 300 K	2060		444
Polycrystalline gold	-	0.01-0.8 L @ 4 K	2125-2110	metallic gold	82
Au/SiO2		10 ⁻³ -700 torr @ 300 K	2129-2106		368
Au (332)	-	0.5 % ML CO-10⁵mbar @ 92 K	2124-2110	metallic gold	83
Au/Na-Y zeolite		0.5 torr @ 77K	2120 2188 2176	Au ^o Au ⁺ CO condensed in zeolite pores	450
Au/TiO2		38 torr @ 300 K	2113-2105 2071-2038 1670 1580	Au ^o bidentate carbonate carboxylate	403
Au/Al ₂ O ₃ /Mo(110)	0.5 ML	10 ⁻⁵ mbar @ 100K	2100	non-metallic gold	367
	20 ML		2110	metallic gold	
Au/H-Y zeolite	1-4 nm	50 torr, 300 K	2111 2138 2169	Au ⁰ Au ⁺ Au ⁺	373
Au/Fe ₂ O ₃		8 mbar CO, 8 mbar O ₂ @ 300 K	2116 2138 2159 2340	Au ⁰ Au neighbouring O Au ⁺ CO ₂	378
Au/TiO ₂	7.5 nm	15 mbar CO, 15 mbar O2 @ 300 K	2116 2151 2187 2351	Au ⁰ Au ⁺ Ti ⁴⁺ CO ₂	376
Au ₈ /MgO/Mo(100)	~6Å	90 K	2102	partially anionic Au	247
Au/TiO2		2.66 kPa @ 298 K	2119 2188 2136	Au ⁰ Ti ⁴⁺ Au, interacting with support	375
Au/Fe ₂ O ₃	2.9 nm	2.66 kPa @ 298 K	2170 2137	Fe ³⁺ Au, interacting with support	377
Au/TiO2	2 nm	5 mbar CO in 1 atm of He @ 300 K	2115 2180 2350	Au ⁰ Ti ⁴⁺ CO ₂	470
Au/Fe ₂ O ₃	4.8 nm 11 nm	100 torr @ 300 K	2107 2123		422
Au/Fe ₂ O ₃	6.5 nm	0.05-1.0 kPa @ 353 K	2111-2104	Au ^o	22
Au/Na-ZSM-5		60 torr CO, 5 torr H ₂ O @ 323 K	2128 2192 1630	Au ^⁰ Au⁺ HCOO ⁻ on Au ⁺	543
Au/Al ₂ O ₃	4 nm	266 Pa @150 K	2108 2154 2186	Au ⁰ oxidized Au Lewis acid site Al₂O₃	541
Au(110)	-	0.1-100 torr @ 300 K	2110 2141	Metallic gold Physisorbed CO	92
Au/ZrO ₂	4.1 nm	0.05-2.5 mbar @ 90 K	2116-2100 2169 2181 2091 2155	Au step on particle support cations support cations Au particle edge Au interacting with OH groups	374
Au/TiO2	4.1 nm	0.009-2.5 mbar @ 90 K	2109-2100 2163 2177 2087 2155	step on particle support cations Ti ⁴⁺ Au particle edge Au interacting with OH groups	374
Au/Al ₂ O ₃ /NiAl(110)	10-600 atoms	0.05-1.0 L @ 90 K	2110-2098		369
Au/TiO2	2-3 nm	1 kPa O2, 1 kPa CO, 1 atm N2 @ 80°C	2112 2362 2335 1690, 1586 1440 1400-1350	Au^0 CO_2 CO_2 bidentate carbonate on TiO ₂ monodentate carbonate formate	400
Au/FeO(111)/Pt(111)	3 nm x 3 Å, 7 x 2 nm	0.1-1.5 L @ 90 K	2108 2131 2165	Metallic gold Au clusters Au ⁺	370, 371
Au/TiO₂/Mo(110)	1.8-3.1 nm	10 ⁻⁷ -10 ⁻² mbar @ 200 K	2126-2118	Au ⁰ , interacting with support	372
Au(110)		10 ⁻⁸ -10 ⁻⁴ mbar @ 110-250 K	2118-2108	Au ^o	90

systems may arise from the specific interaction of gold with oxygen rather than CO. However, the finding that annealing strongly attenuates size effects indicates that the nature of the support and its defect structure may be critical for nucleation, growth and stabilization of very small Au clusters upon which CO adsorbs more strongly. Just as we have observed slight differences between forms of iron oxide, Okumura *et al* had previously observed differences in the annealing behavior of gold on different forms of TiO_2 and found that defect structure of the support strongly controlled the particle size distribution after calcination and therefore the reactivity (385).

Particle size effects have been observed in the adsorption of other molecules as well. Sykes *et al* observed that styrene was bound much more strongly to small gold particles on titania as compared to a thick gold film as the desorption temperature was shifted from 350 K to 410 K (278). Furthermore, the desorption signal from the titania surface itself at 250 K was lost when depositing only 0.25 ML gold on the titania surface. This result implies that during the TPD experiment, styrene on the titania surface was mobilized and migrated to the gold particles where it then adsorbed in a much stronger manner.

It has been speculated that molecular oxygen could be adsorbed (and possibly dissociated) on small gold particles but to date no direct observation has been made. Zhu *et al* report the adsorption of molecular oxygen on gold particles on HOPG using XPS at room temperature following exposure of oxygen at 1 atm (386). However, as the particles are quite large (20-50 nm) it seems unlikely that they possess adsorption behavior greatly different from bulk gold, which has been conclusively shown to be inert to such exposures (95) and therefore it seems exceedingly unlikely that molecular oxygen could possibly be a stable species in UHV at 300 K on such structures.

Choi *et al.* showed desorption of oxygen from Au/SiO₂ following large (on the order of 10,000 L of gas) exposures of NO₂ or O₂ (387). Oxygen was observed to desorb in a peak at about 800 K. The authors speculated that this peak arises from desorption of sub-surface oxygen. The amount of oxygen was shown to increase as the exposure temperature of the sample increased, indicating the chemisorption process is highly activated. The authors concluded that an interaction between the gold and the silica support was responsible for the increased chemisorption of oxygen on the supported gold surface (as opposed to gold single crystals).

In a similar fashion to the work on single crystals, the adsorption of atomic oxygen on gold particles on titania has been studied by Campbell and co-workers using a hot filament to dissociate oxygen (196). Campbell's group observed a very strong shift to higher oxygen desorption temperatures from 530 K to 645 K as the amount of gold

Figure 22

Turnover frequencies from (a) $Au/TiO_2/Mo(110)$ model catalysts grown in UHV (b) Au/TiO_2 catalysts prepared by deposition precipitation for CO oxidation at 300 K as a function of the Au particle size. (232 (data from 192, 395))

deposited was decreased from 6 ML to 2.3 ML. For the thinnest islands, estimated to be a single monolayer, no oxygen desorption was observed below 710 K even though LEIS measurements of the attenuation of the gold signal confirmed the presence of oxygen on the surface. Due to their stronger binding of atomic oxygen, the authors suggested that gold particles below a critical thickness may have greatly enhanced oxygen dissociation capability. In addition, CO oxidation activity was demonstrated by the evolution of CO₂ from the room temperature sample surface upon exposure of the oxygen covered Au/TiO₂ model catalyst to 10⁻⁶ torr of CO (388). TPD experiments showed total depletion of oxygen from the surface following CO exposure. Rate constant estimates based on the rate of disappearance of oxygen revealed that particles of two-layer thickness were approximately twice as reactive as one layer thick particles. As seen in the first experiments by Campbell's group, the thinner particles bind oxygen more strongly so it is not as easily available for reaction. At the same time, the thinner particles may have a diminished CO adsorption capacity. An apparent activation energy of 11.4 kJ/mol was reported for the two-layer thick particles, which is not remarkably different from results reported by Outka and Madix on Au(110) or by Haruta for real catalysts.

Martra *et al* have recently observed that smoother gold particles on SiO_2 were more reactive for CO oxidation although their CO adsorption capability was reduced (389). Like Bondzie *et al* (388), the authors theorized that oxygen was bound too strongly for reaction on particles with a higher number of low coordinated atoms.

The production of CO_2 for temperatures as low as 65 K has been observed by Kim *et al* when impinging a molecular

beam of CO on a Au/TiO₂(110) sample that had previously been exposed to atomic oxygen (390). The authors reported that CO₂ production is maximized for an oxygen coverage of about 0.1 ML. Although site blocking cannot be ruled out, the authors speculated that electronic effects were likely responsible for the reduction in CO adsorption and consequent loss of reactivity at higher oxygen coverages. Significantly, no particle size effect was observed, as the amount of CO₂ increased with the amount of gold deposited (in the absence of gold no CO₂ is produced). This result combined with findings from the Campbell group (388) suggests that mechanisms involving atomic oxygen are only slightly particle size dependent. Furthermore, the tendency of larger particles to be more reactive is counter to observations made when the oxygen is not atomically dosed and seems to imply that oxygen dissociation is the ratelimiting step. Mullins' group also reported that for temperatures above the desorption temperature of CO, the reaction rate decreased rapidly following exposure to CO, but then rose slightly showing a second maximum in production, frequently several seconds later (391). The authors have speculated that two different mechanisms (possibly one E-R mechanism and one L-H mechanism) must be present in order to explain the presence of the second maximum.

Recently, Mullins' group has found evidence for a molecular oxygen species on Au/TiO_2 model catalysts. Some molecular oxygen was found to be delivered to the surface as a by-product of the oxygen atom source used in the experiments described above (392), and is likely in some electronically excited state. The reactivity of the molecular oxygen species as compared to atomic oxygen is the subject of further research (393).

Goodman's group (192,394) made use of a high pressure cell (40 torr) and were able to observe CO₂ production by a planar Au/TiO₂ catalyst when exposing the surface to a 1:5 CO:O₂ mixture at room temperature. They observed a strong particle size effect with those particles in the range of 2.5-3.5 nm showing a maximum in activity. Although Haruta originally observed that the activity increases continuously as the particle size decreases (9,187,343), a careful examination of the data presented by Bamwenda et al (395) reveals extraordinary agreement with the findings of the Goodman group (192,394). As discussed above, Valden et al related this maximum in activity to the transition between metallic and non-metallic particles. Variations in particle size also gave variations in the apparent activation energy (3.5-6.0 kcal/mol). The authors concluded that this must result from an intrinsic property of the particles and not from any geometric effect due to the changing ratio of perimeter atoms to surface atoms. Gold particles with an average diameter of 2.4 nm were shown to lose activity rapidly over the first hour of exposure to the gas mixture at room

Figure 23

Temperature programmed reaction spectra for CO_2 production following CO and O_2 exposure at 90 K to Au_8 and Au_{20} clusters on MgO. (405)

temperature and AES showed an increase in carbon on the surface. Efforts to regenerate the catalyst with oxygen were not successful in spite of the removal of carbon.

Valden *et al* (192,394) and work by Haruta (9, 191, 395, 396) report a strong size effect: particles above 5 nm exhibit virtually no activity while particles below 5 nm show good activity for room temperature CO oxidation, exhibiting a maximum in activity at 2.8-3.2 nm as shown in Figure 22. The observation of such size effects in catalysis is not new (397,398). However, the difference in activity between bulk gold and supported gold particles is rather dramatic. In the widely studied oxidation of CO, oxide supported catalysts have been shown to be at least two orders of magnitude more active than unsupported gold (399). However, how and why supported gold particles experience an increased reactivity is still largely unknown.

However, it appears likely that the exact size range which gives a maximum in activity may depend upon the reaction or the support or even the catalyst preparation method. For example, Schumacher et al. have recently reported that the maximum in activity below 2 nm for their Au/TiO₂ catalyst (400). Prati and co-workers have found for the oxidation of ethylene glycol to glycolate that the activity increased with decreasing particle size for Au/Al₂O₃ and Au/TiO₂ catalysts but that the rate exhibited a maximum at some intermediate size for activated carbon supported gold catalysts (401,402).

Figure 24

(a) Front and (b) side views of the CO oxidation reaction on an Au₈ cluster. CO is conceived to adsorb on-top of the cluster while oxygen is conceived to adsorb molecularly at the particle periphery. (247)

Some authors have even reported that catalytic activity and particle size had little or no relationship (403). The picture is often muddled by the lack of a monodisperse size distribution. Heiz and co-workers have attempted to address this problem by using a mass spectrometry size selection technique, by which clusters from 1-20 gold atoms in size were examined on an MgO thin film substrate (247,404-406). Sanchez et al found that the onset of CO oxidation activity occurred at 8 atoms, below which no activity was observed and above which catalytic activity increased in an oscillatory fashion. The CO₂ desorption temperature was also found to be cluster size dependent. As shown in Figure 23, for Au₈, CO₂ was produced in two peaks, a low temperature feature at 140 K and a larger feature at 240 K. As the cluster size increased, the desorption temperature of the high temperature feature also increased (up to 500 K for Au₂₀) while the low temperature feature remained constant. It is notable that CO was observed to desorb between 150 and 180 K from the Au₈ clusters in the absence of oxygen. Therefore, as has been seen in gas phase experiments, the adsorption of CO and O_2 must be cooperative such that CO (or at least some kind of transition state) is stabilized to much higher temperatures in the presence of oxygen. Similarly, oxygen may be stabilized by the presence of CO, although at this point the adsorption/desorption behavior of oxygen on Au₈ remains a mystery. Supporting calculations indicated that although the dissociation of oxygen is thermodynamically favored, the large barrier to reaction implied that the reaction should proceed through molecular oxygen.

Calculations also showed that while the adsorption of oxygen was highly dependent upon the cluster size, the adsorption of CO is relatively cluster size insensitive. The reaction was conceived to occur with the adsorption of CO on the cluster and the adsorption of oxygen on the particle periphery as shown in Figure 24. CO₂ production was greatly enhanced for Au clusters on defect rich films as compared to

Figure 25

The Au 4f region of XPS spectra obtained for a Au₅₅ clusters deposited on a silica wafer and Au particles formed by metal evaporation on the same support following oxygen plasma treatment. The results show a high oxidation resistance of the closed-shell Au₅₅ cluster. (418)

clusters on defect poor films. Likewise, simulations indicated that the reaction barrier of 0.8 eV was lowered to 0.5 eV on Au₈ clusters above defects due to the 0.5 eV charge that is predicted to spread over the 8 atom cluster (247). Interestingly, Pd₈ clusters did not exhibit the same defect dependent reactivity (405). Doping the clusters with one strontium atom was found to increase the reactivity of the clusters and clusters as small as Au₃Sr showed appreciable activity (407). Perhaps surprisingly, the CO desorption temperature was not greatly different from that observed by Hollins and Ruggerio for CO on Au(332) (83). Moreover, IR data reveal a single peak at 2102 cm^{-1} , for the C-O stretching frequency, which is in agreement with other data for CO on neutral bulk gold surfaces (see Table 2). This would seem to indicate an absence of special binding sites connected to anionic particles and the origin of the enhancement of CO adsorption on Au₈ clusters remains unclear. For the Au₈ clusters, the oxygen exposure-CO exposure- TPR cycle (to 350 K) could be repeated several times without observing any drop off in CO2 production. This stability implies that the dispersed clusters were strongly enough bound to the MgO defects that they did not sinter.

Both Goodman's group (408) and Gates' group (409) have succeeded in preparing Au₆ clusters using more conventional catalyst preparations. Goodman and co-workers employed a

Au₆(PPh₃)₆(BF₄)₂ precursor on a TiO₂(110) support which after electron stimulated desorption of the ligands gave highly dispersed Au₆ clusters as seen by STM (408). In order to avoid problems with phosphorus contamination, Gates and Guzman opted to use Au(CH₃)₂(C₅H₇O₂) on MgO powder (409). After holding the catalyst in a He atmosphere at 373 K for 2 hours, EXAFS spectra indicated the formation of Au₆ clusters. However, the reactivity of these novel systems has not been established.

Based on the above results and results from gas phase experiments, some researchers have advanced the idea of "magic number" clusters (410) that exhibit extraordinary activity. From their combined analysis of activity data, TEM and Debye function analysis (DFA) of the XRD spectra, Cunningham et al proposed that icosahedral Au₁₃ particles on a Mg(OH)₂ support acted as the active species (411,412). The 13 atom clusters were conceived to be essentially constructed entirely from 5-fold coordinated atoms. Calcination at higher temperatures led to applomeration and thereby less active catalysts, while calcination at lower temperatures left oxidic gold oligomers on the surface that failed to show appreciable activity. Examination of a used catalyst showed a complete absence of icosahedral structures and truncated decahedra had formed instead. In a comparison of supports, however, DFA identified fcc cubooctahedral gold structures from 1-8 nm to be the catalytically active species on TiO_2 (413). For example, Au₁₄₇ cubooctahedral particles formed the dominant species for a 1.3 wt% Au/TiO₂ catalyst. Therefore, Cunningham and co-workers concluded that the catalytically active structures are dependent on the support and must be considered on a case-by-case basis.

Cleveland *et al* found in their examination of gold particles grown in solution that certain size clusters were preferentially formed as determined by XRD (210). The "magic sizes" of 75, 101, 146 atom clusters were found to be remarkably stable as XRD patterns of the same samples stored in air taken one year later were identical.

One of the first examples of a magic cluster is Au_{55} which was discovered by Schmid *et al* (414) and has since seen extensive study (415-417). Boyen *et al* recently confirmed that Au_{55} clusters display unusual chemical behaviour (418). They deposited two types of gold nanoparticles on a silicon wafer, Au_{55} clusters and non-size selected particles ranging in average size from 1.0 nm to 8.0 nm. XPS measurements were used to determine the degree of oxidation after exposure to an oxygen plasma. For the larger particles, the XPS spectra revealed four peaks for the Au 4*f* region, including two associated with Au_2O_3 produced as a result of the plasma and two associated with metallic gold in the particle interior. Remarkably, the particles in the region around 1.6-1.3 nm did not show significant oxidation as shown in Figure 25. However, particles under 1 nm exhibited significant oxidation, seemingly indicating a maximum in oxidation resistance for the Au_{55} clusters. The valence band region spectra showed that the Au_{55} clusters showed small intensity right up to the Fermi level as is seen for bulk gold. Therefore the authors concluded that the remarkable resistance to oxygen was not a consequence of electron confinement but rather due to an electronic closed shell configuration.

6 Connection to real catalysts

6.1 What is the active site?

Currently, the real nature of gold catalysis, particularly with regard to the well-studied low temperature CO oxidation reaction, remains a subject of much debate. In general, as depicted in Figure 26, CO is envisioned to adsorb on the gold particles themselves, probably on edge or step sites (3,4). The adsorption site of oxygen is less clear, although mechanisms involving oxygen adsorption on the support or at the particle-support interface are often invoked as will be discussed in Section 6.2. The following ideas or combinations thereof have been proposed by researchers to explain the phenomenal reactivity of gold nanoparticles:

- 1 Small gold particles possess unique adsorption sites that allow for such reactions as the adsorption or dissociation of oxygen due to the presence of highly uncoordinated atoms, (190,197,221,222,369,419,435,436).
- 2 Small gold particles have unique electronic and thereby chemical properties as a consequence of the quantum size effect. Some have amended this idea with the corollary that the thickness of the gold particle is the critical parameter. The role of the support is merely to stabilize these small particles (192,196,323,388,392).
- 3 The active site for gold catalysis involves Au^* whose presence may result from the catalyst precursor or the calcination treatment (378,444-458,460).
- 4 The active site for gold catalysis is an anionic gold (Au') species which results as a consequence of the interaction of a particle with a F center defect in the underlying oxide support and which allows for the adsorption (and possible dissociation) of oxygen as shown in gas phase experiments (170,184,185,220,247,374,462-465).
- 5 The active site for catalysis involves the interface between the particle and support. For CO oxidation it is generally proposed that CO adsorbs on the surface of the gold particle whereas oxygen adsorbs on the metal oxide support and moves to the metal/support interface or adsorbs there directly (226,365,403,470-472,474,475).

We have previously discussed size effects in gold catalysis, which arise either from electronic effects, structural effects or a combination thereof. In spite of the data found

Figure 26 Schematic representation for CO oxidation pathways over Au/TiO₂. (3)

in both the catalytic and surface science communities, particularly the dramatic results from size selected clusters, many experiments involving real catalysts report that size effects are not a critical factor in determining activity (365,403,419,420). We are therefore left to question what the controlling factors for CO oxidation really are.

Recently Haruta has established a group of conditions whereby active gold catalysts can be made (3,4). First and foremost Haruta asserts strong contact is necessary between gold and the underlying support. Although calcinations at high temperatures generally leads to agglomeration of particles which is assumed to be undesirable (395,421-423), Haruta and co-workers found an increase in activity with increasing calcination temperature between mechanically mixed Au colloid/TiO₂ powder samples calcined at 573 K and 873 K (424). TEM results showed the more active catalyst, in spite of its larger particle size, had better contact with the support. As a second point, Haruta argues that the proper support must be chosen. Basic supports are preferred for the type of deposition precipitation method that has been shown to be effective for the preparation of active catalysts. Traditional impregnation methods with gold chloride are unsatisfactory because during calcination of the catalyst, the gold particles experience severe agglomeration as HAuCl₄ interacts only weakly with the support. The deposition precipitation method favored by Haruta (although many other methods have achieved at least limited success as well) relies upon an exchange between gold chloride and NaOH such that Au(OH)3 is formed and precipitates on the support. The dispersion and size of the particles, and therefore their catalytic activity, depend critically upon the pH used (typically 6-10) and the amount of gold in solution (423,425-432). After washing to remove Cl and Na ions, the catalyst is dried and calcined. In contrast to HAuCl₄, AuOH₃ applomerates only minimally under calcining conditions. However, with the deposition precipitation method supports such as activated carbon (433) and zeolites (434) cannot be used due to their high acidity (isoelectric point). (It is interesting to point out that graphite substrates have often been used in photoemission experiments designed to explore the electronic properties of gold nanoparticles and that such particles appear to share the similar electronic character to gold particles on oxides). Finally, as a third condition, particles should be in a particular size range, although under the right circumstances particles as large as 10 nm can exhibit high activity. However, in the case of real catalysts, size distributions tend to be rather wide and difficulties may be encountered in the detection of very small particles, which could be the active species.

The effect of geometry has been discussed from a theoretical standpoint, but it is important to mention that some recent experimental results from real catalytic systems have also emphasized the influence of geometrical effects. As mentioned earlier Baiker's group found that the activity of Au/TiO₂ was much higher than Au/ZrO₂ found catalysts with the same particle size 359,419. Originally ascribed to support effects, HREM revealed that the differences in activity may arise from differences on particle shape. Au/TiO₂ catalysts had more highly uncoordinated atoms and were thereby more active as has been predicted by Nørskov and coworkers 190,222.

Mohr et al have identified reaction sites on the corners of gold particles for hydrogenation of acrolein to allyl alcohol (435,436). In a comparison of Au/TiO₂ and Au/ZrO₂, Mohr et al attributed differences in reactivity to the degree of rounding of the particles. Although, particles on the two supports were nearly the same size, Au/TiO₂ was approximately twice as active and more selective as well for the production of allyl alcohol as Au/ZrO₂. From an analysis of TEM images, the authors attributed these differences in reactivity to the presence of more highly uncoordinated atoms on particles supported by titania. More recently, by HREM it was found that indium wets the flat surfaces on gold leaving only the edges exposed (436) as shown in Figure 27. The authors found that zinc oxide supported 10 nm gold particles covered with indium were still half as reactive as those not covered by indium with a much better selectivity to allyl alcohol. This led the authors to conclude that the exposed corners and edges must be the reactive site for the selective hydrogenation of the C=O group as ZnO itself is not active for the reaction.

As mentioned above, Campbell (196) and Goodman (192) have suggested that two-dimensional islands of gold may exhibit unusual activity as compared to their 3-D counterparts. However, in our own work we failed to see any differences in CO adsorption between gold monolayers on FeO as compared to larger 3-D particles (370). Boccuzzi et al has previously suggested that 2-D rafts of gold may easily absorb negative charge from the support based on their observation of bands around 2050 cm⁻¹ for CO adsorption on reduced Au/TiO₂ (437). As mentioned previously, Blick et al observed the presence of two-dimensional particles of gold by TEM and Mössbauer spectroscopy. The authors found that for the oxidation of methane, 2-D rafts were essentially inactive and served only to block methane coupling reactions that probably occur on oxygen vacancies now covered by gold (299). The authors proposed that 3-D particles of gold were necessary for reaction and that the reaction took place at the particle support interface. Although the presence of Au-" sites associated with two-dimensional gold rafts was observed by CO adsorption on both Au/TiO₂ and Au/Fe₂O₃, Boccuzzi et al found that the reverse water gas-shift reaction were only catalyzed by three-dimensional gold particles as the resulting CO formed was found only on Au^o sites (438).

Early EXAFS and XPS results detected the presence of oxidic gold in active catalysts (439-443). Lee and Schwank were among the first to suggest that such cationic species acted as the active sites in gold catalysis (444). Based on IR data which showed CO adsorption 2040 cm⁻¹, considered to be on sites other than metallic gold, and considering the lack

Figure 27

High resolution TEM images of two different indium covered Au particles (left side) on ZnO with corresponding assignment of structures (right). (436)

of activity for unsupported bulk gold, the authors proposed that these adsorption sites may be related to gold that is electronically modified by the support due to charge transfer. More recently, infrared spectroscopy of CO adsorption on gold catalysts has revealed features between 2150-2160 cm⁻¹ (as seen in Table 2) which has led some researchers to invoke a mechanism which involves Au⁺ (378,382,445). XPS and FTIR experiments by Galvagno and co-workers have shown a loss of Au⁺ species over time when Au/Fe₂O₃ catalysts were exposed to reactive mixtures (378,446). The authors concluded that the deactivation of the catalyst over time was directly related to the loss of Au⁺ and the growth of Au⁰. In most studies, an increase in the calcination temperature resulted in a decrease in the catalyst activity. Park and Lee tied this effect to the presence or absence of oxidic gold (447). As calcination temperature of Au/TiO₂ was increased from 373 K to 673 K, the activity was observed to fall by a factor of 10. At the same time, XPS and XANES data showed a transition from an oxidic gold species similar to Au(OH)₃ to a metallic gold species whose photoemission spectra were indistinguishable from a gold foil. Hao and co-workers later confirmed this result for Au/Fe₂O₃ catalysts (448). Hao et al speculated that the calcinations process transforms Au³⁺ to Au^{d+} essentially giving Au a similar electronic structure to Pt. In similar work using Mössbauer spectroscopy, Kobayashi et al found that the most active Au/Mg(OH)₂ catalysts also had the largest amount of Au¹⁺ (449) and thereby concluded that oxidic gold is the active species for CO oxidation. Ichikawa and co-workers also asserted that Au^+ is the active component based on their finding that Au^{+/}/NaY zeolite catalysts were considerably more active than their Au^o counterparts (450) for reduction of NO. The authors proposed that positively charged gold species acts as a good catalyst due to its unfilled *d*-band (451). The active site was conceived to be Au¹⁺ ions that exist at the interface between metallic gold particles and the zeolite support (450). Furthermore it was found that Au¹⁺ catalysts were more active than Au^{3+} (452).

Unlike the findings from Ichikawa's group, other investigations have found that higher oxidation states were even better catalysts. Waters *et al* saw, in their comparison of Au on various supports, that the activity for methane oxidation increased directly with the oxidation state of gold as observed by XPS (453). The authors hypothesized that oxidized gold had a greater tendency to chemisorb oxygen. Gates and Guzman recently determined that Au³⁺ is the active species for hydrogenation of ethylene by gold (454). XANES data revealed a direct correlation between the coordination number and the activity of MgO supported gold catalysts. An increase in activity was also observed with increasing amounts of Au³⁺ and the best catalyst was found to be a mononuclear gold complex formed by deposition of

 $Au^{3*}(CH_3)_2(C_5H_7O_2)$. Similar to Ichikawa's group, Wan and coworkers ascribed deactivation of their Au/Y zeolite catalysts to transformation of Au^{3*} to Au^0 as seen by XPS (455).

Hutchings and co-workers found that uncalcined iron oxide supported gold catalysts were considered more active than those calcined at 400°C (456,457). They attributed this large difference in reactivity to the presence of goldoxyhydroxide (AuOOH.H₂O) as observed by Au¹⁹⁷ Mössbauer spectroscopy. Unlike many other studies, the activity of their catalyst increased over time. This could be an indication that further changes to the catalyst (such as partial reduction by CO) are beneficial to the catalyst's performance. TEM examination showed that while the relatively inactive calcined catalysts showed particles in the 3-5 nm range, which are generally thought to be of ideal size, the more active dried samples revealed a more homogeneous distribution of gold. Kang and Wan found that iron modified Au/Y zeolite catalysts could be regenerated with water vapor (458). Based on this result, they proposed that gold hydroxide in intimate contact with the oxide support acted as the active species in CO oxidation.

Some have even proposed that gold is essentially incorporated into the support itself (365,455,459). If gold possesses the same valence as the support cation, then it is reasonable to suggest that gold could simply replace the support metal. Sze *et al* found evidence of this behavior using Mössbauer spectroscopy for Au/Fe₂O₃ catalysts (459). Using a co-precipitation preparation method it was possible to insert gold into the iron oxide lattice. However, upon annealing above 300°C, gold was once again segregated from the Fe₂O₃ phase. Wan and Kang had similarly proposed that gold was incorporated into the lattice of a Y zeolite during preparation and that in fact the point at which this substitution occurred then acted as a nucleation site for further gold deposition (455).

Flytzani-Stephanopoulos and co-workers have recently demonstrated that gold nanoparticles are not the active species for the water-gas shift reaction seen on Au/CeO₂ catalysts (460). The authors found that using a leaching process almost all the metallic gold could be removed from Au/CeO₂ that had been prepared by deposition precipitation or co-precipitation, leaving behind a highly dispersed gold species. These leached catalysts exhibited comparable activity to the as prepared catalysts leading the authors to conclude that metallic gold must not contribute to the catalyst's reactivity. Flytzani-Stephanopoulos and co-workers support a model in which Au is integrated into the ceria support. The authors suggest that the role of Au is to increase the amount of surface oxygen in ceria (461).

In contrast to the situation with cationic gold, relatively less attention has been focused on the presence of anionic gold as the active species. However, as mentioned above, oxygen vacancies on titania and magnesia are preferable nucleation sites for gold (246,251-255). In addition, Heiz and co-workers have demonstrated that gold above such defects will exhibit higher activity (247). Calculations indicate that gold above the oxygen vacancy is electron rich as a charge transfer is imagined to take place from the support to the gold particle. Such an active site would also be consistent with gas phase experiments which demonstrated that oxygen preferred to adsorb on negatively charged clusters (170). Since there is some evidence that oxygen adsorption (and dissociation) is the rate-limiting step (87,196,388), the presence of anionic clusters is perhaps crucial to the reaction's success.

Claus and co-workers, in their examination of acrolein hydrogenation on Au/TiO₂ and Au/ZrO₂ catalysts, concluded that anionic gold and the support acted together to produce the active site for allyl alcohol production (323,462). As in Heiz's model, electrons trapped at paramagnetic F-centers (oxygen vacancies) can be transferred to the gold particles. The electron rich gold particles may then have heightened interactions with various adsorbates. Correlating XAFS data with catalyst activity, Kageyama et al found that the most active catalysts also had the highest degree of Ti-Au coordination (463). It was proposed that Au species were anchored in the surface of TiO₂ and that the higher catalytic activity may be the result of charge transfer from Ti to give negatively charge gold particles. Manzolli et al, in their study comparing Au/TiO₂ and Au/ZrO₂, concluded that the higher activity of the titania supported catalyst resulted from the presence of oxygen vacancies which donate electrons to small gold particles (374).

Based on a combination of XPS and reactivity data, Horvath *et al* proposed a mechanism whereby gold supported on iron oxide which is originally cationic becomes neutral or even anionic after oxidation due to a charge transfer from the support to the gold particle (464). TEM data showed that the particle size distribution was essentially unchanged after oxidation indicating that the binding energy shift observed probably arose from a change in the chemical relationship between the gold and the support. Andreeva and co-workers have also invoked the idea of charge transfer from the support to the gold particles to explain the strong synergistic effects observed in the oxidation of benzene over $Au/V_xO_y/TiO_2$ catalysts (465).

Nieuwenhuys' group observed that while preoxidized samples of Au/TiO_2 were initially inactive, their activity increased following exposure to CO and O_2 (376). On the other hand, prereduced samples exhibited a high activity from the reaction onset. However, extensive reduction of catalysts has been shown to be detrimental to their activity in many cases (423,466-468).

The presence of special sites created by the metal support interface is a common explanation invoked in the theory of catalytic mechanisms (469). Since neither bulk gold nor bulk

titania are active (or are at least significantly less active than Au/TiO₂) for low temperature CO oxidation (403), some have theorized that special sites created by the interface must play a key role in the catalysis (226,403,430,470-472). Strong evidence for such sites has been presented by Vannice and co-workers by their use of inverse catalysts of titania particles deposited on gold substrates. When depositing TiO_x on gold powder, appreciable activity was observed (403), leading the authors to conclude that activity has no relationship to quantum size effects, but rather is primarily a function of the metal-support interface. Efforts to identify specific adsorption sites for CO on Au/TiO2 that arise from the interface proved inconclusive using TPD (471). This phenomena is not unique to Au/TiO₂, as Fu et al have seen similar synergism in Au/CeO2 catalysts (365). From their results, the size of the gold particles did not appear to play a role in the catalyst's activity. However, the size of the CeO₂ crystals was a critical parameter. Guczi and co-workers have proposed a mechanism whereby there must be contact between gold and the active oxide phase based on their study of various Au/FeOx catalysts (472). They speculated that the interface between amorphous iron oxide and gold is electron rich, resulting in an ideal place for CO adsorption and oxidation. However, in our own work, we also failed to see unique adsorption sites for CO that must be ascribed to the metal-support interface (302,370). However, FTIR experiments of CO adsorption have shown that on many surfaces adsorption features are seen at 2090-2105 cm⁻¹, which are generally outside the normal range for metallic gold (2105-2125 cm⁻¹) (473). Therefore these features have been assigned to particle edges at the support interface. It remains unclear, however, how these adsorption sites are related to the unique catalysis by gold.

Recent calculations by Hammer and Molina for CO oxidation by a 34 atom gold cluster on MgO revealed that the support serves to assist in the stabilization of the reaction intermediate (474). Various types of interfaces were tested for adsorption of oxygen and CO and for their reaction as depicted in Fig 28 a-c. CO was found to bond to the gold cluster at a low coordinated site near the edge. CO adsorption at the interface of between the support and clusters with a high degree of wetting (as in 28a) was found to be disadvantageous due to steric effects as the CO was repulsed by the support. Like Sanchez et al (247), a mechanism involving molecular oxygen was favored since the barrier to dissociation of oxygen is too great (more than 1 eV). Oxygen was found to bond to both the cluster and support, ultimately forming a CO-O-O complex at the cluster's edge with the most favorable arrangement shown in Figure 28b. When the complex rearranges as depicted in Fig. 28d, then the barrier to reaction is lowered significantly and CO₂ is readily formed. The second oxygen remaining on the gold particle is even

more easily reacted with CO. Fig. 28e shows an alternate arrangement of the reaction at the interface. However, in this case the adsorbates are not bound to lowly coordinated gold as in 28b or d so the reaction is less favorable.

Ajo et al have shown that propylene may have unique adsorption sites on the particle-support interface (475). For gold particles with an estimated average thickness of 1.5 ML on TiO₂(110), two distinct peaks (150 K, 240 K) were seen from propene desorption at low coverages. The low temperature peak was assigned to propene desorption from bulk gold surfaces (as seen by Davis and Goodman (104)). As coverage was increased, a third peak emerged at 190 K which corresponded to propene desorption from titania. Ajo et al speculated that the high temperature feature resulted from sites on the particle-support interface. At low coverages, propene was adsorbed directly onto the particles and was effectively trapped there by propene that adsorbed on the titania and then migrated to the interface. Thicker gold islands on TiO₂(110) exhibited similar behavior, although the loss of the high temperature tail to the TPD spectra was observed. The authors ascribed this high temperature tail to propene desorption from the interface of the support with one monolayer thick particles, which were no longer present for higher gold loadings. It was therefore estimated that the desorption energy from such sites was about 20% higher than that from edge sites of bulk gold particles.

Schüth and co-workers have employed a novel solution technique to develop highly active gold particles that have limited contact with the support (476). The catalyst was prepared by depositing $AuCl_4$ on an alkylammonium functionalized silica support. As the support is essentially covered with an organic, the metal-support interaction should be negligible and therefore, the authors reject hypotheses that involve special sites created by the metal support

interface. Noting that active catalysts have been created on a variety of supports, they assert that gold particles must only be sufficiently small in order to display good activity.

As mentioned previously, the adsorption site of oxygen is largely a mystery. Oxygen is known to adsorb readily on oxygen vacancies on metal oxide supports such as titania (477,478). However, oxygen does not merely fill vacancies and as both molecular and atomic oxygen can coexist on the titania surface under the right circumstances (479), the determination of the reactive form of oxygen is complex (as discussed below). As indicated in Fig 26, Haruta and co-workers have described mechanisms whereby oxygen is adsorbed on the support (either dissociatively or molecularly) and then is transported to the particle-support interface where it reacts with CO. Using FTIR, Magkoev et al observed that for gold deposited on a highly reduced TiO_x support the rate of disappearance of CO was far greater than for Au/TiO₂ upon exposure to oxygen (480). The authors rationalized this by proposing a mechanism whereby the presence of oxygen vacancies greatly facilitated the adsorption of oxygen and subsequent reaction. Complementary XPS experiments revealed that the Au/TiO₂ catalytic surface could be reduced following prolonged exposures to CO at 500 K. Since reduction did not occur in the absence of gold, the authors concluded the presence of the Au/TiO₂ interface must act as the active site for catalysis. However, from their study comparing the production of CO₂ from Au/TiO₂, Au powder and TiO₂ powder, lizuka et al observed that oxidized Au powder produced about 200 times the amount of CO_2 as oxidized TiO₂ powder after exposure to CO (481). Therefore since it appeared that oxygen reacts only with great difficulty on $\text{TiO}_{2},$ the authors concluded that oxygen activation (whether dissociative or not) must take place on the surface of gold particles or at the particle-support interface.

Another factor to consider of course with regard to the activity of gold concerns the presence of impurities. It has been shown, in numerous examples (482-484), that bimetallic particles involving gold are superior to gold alone. Therefore, it stands to reason that impurities present in gold actually have a promoting effect in catalysis. In fact, as previously discussed, such impurities have been found to facilitate oxygen adsorption and dissociation (96-100). Iizuka and co-workers recently presented evidence that the reactivity of Au powder could be tied directly to the concentration of silver at the surface (485). Although, gold possesses a very low surface energy, impurities may be stabilized at the surface in a reactive environment and thereby contribute significantly to catalysis (486).

It should be clear that from the above discussion that different reactions may involve different sites and that the principles controlling the chemistry for any particular reaction may not be applicable to others involving very similar catalysts. In other words, the "magic" of gold nanoparticles cannot be simply tied to one particular attribute or active site. General statements regarding the origin of gold's astonishing activity should therefore be avoided as the reactivity for any given reaction is likely a result of a combination of factors. In fact, as will be discussed in the next section, multiple reaction pathways involving multiple active sites may exist even for a simple reaction such as CO oxidation.

6.2 Mechanisms of reaction

Another compelling problem in gold chemistry is an understanding of the catalytic mechanisms. For CO oxidation, there are at least three mechanisms which have received support.

- 1 CO reacts with atomic oxygen
- 2 CO reacts with molecular oxygen
- 3 The reaction proceeds through a complex scheme involving surface hydroxyls.

Of course, as with any scientific problem of importance, the situation is not so simple. As one can see from our discussion of the active site there exist numerous variations and possibilities. Some researchers have even proposed that the mechanism could be different for different types of supports (226,413,487) or that multiple pathways are concurrently active.

Atomic oxygen on the surface of gold single crystals has clearly proven to be capable of oxidizing CO and catalyzing other reactions as discussed in Section 2.3. However, the critical question remains: how does oxygen dissociate on gold surfaces. As mentioned in Section 4, the dissociation of oxygen even on highly stepped surfaces is an extremely endothermic process (197). Therefore, it must be believed that the barrier to dissociation of oxygen on very small supported particles is substantially lowered, perhaps due to electronic effects. As of yet there is no direct evidence that small gold particles dissociate oxygen although, recent calculations of O₂ on a ten atom cluster indicate this may be possible when gold is relatively uncoordinated (221). In addition, the adsorption of atomic oxygen appears to be rather strongly dependent on particle size indicating that the dissociation of oxygen is also likely to be favorably influenced by particle size (196).

Although CO oxidation occurs through dissociated oxygen atoms on other metal surfaces such as Pt (488), mechanisms involving molecular oxygen have been proposed for gold catalysts by many researchers (226,247,375,377). Since CO oxidation has previously been shown to occur with molecular oxygen on silver surfaces (489), it seems logical that gold may also favor such a reaction pathway. However, as molecular oxygen does not adsorb readily either dissociatively or intact on gold surfaces, the mechanism has proven to be difficult to elucidate.

A remarkable feature of the CO oxidation reaction catalyzed by small gold particles is the incredibly low temperatures by which these reactions are possible. For

example, highly dispersed Au/TiO2 catalysts have shown activity down to 209 K (490, for example) and gold clusters have shown reactivity at liquid nitrogen temperatures (169,185). Although it is known that bulk gold is a catalyst for CO oxidation at high temperatures (300°C) (491-493), this reaction may occur via a completely different mechanism than the low temperature reaction (or even catalyzed by impurities). It has been argued that the fact that CO oxidation on small gold particles occurs at low temperatures with very low activation energies implies that it cannot proceed through a mechanism involving a highly energetic process such as oxygen dissociation (220,474). Schubert et al (226) and Iwasawa and co-workers (375,377,494) have reported that $^{36}\text{O}_2$ and $^{32}\text{O}_2$ demonstrated no mixing on the catalyst surface. Furthermore, when reacting C¹⁶O with ³⁶O₂ the only reaction product seen was C¹⁶O¹⁸O with no double labeled CO₂ of either type. These results indicate that lattice oxygen was not participating in the reaction in any fashion and suggest that oxygen did not dissociate readily on the surface. This would seem to argue against any type of Marsvan Krevelen mechanism (454) as has been proposed in some work (487,496,497). Schubert et al proposed a mechanism in which oxygen adsorbed at the metal/support interface reacts with CO through one of two routes (226). In the first route, molecular oxygen adsorbs on the support surface before migrating to the interface where it can dissociate and react with CO through a spillover process. In the second mechanism, a direct reaction between CO and molecular oxygen is suggested to occur through a noncarbonate-like transition state. While Behm's group favors the first of these mechanisms, Iwasawa's group favors the latter (375,377). Hao et al observed O_2^- ions on the surface of Au/ZnO using EPR after exposure to oxygen at room temperature (498). Upon introduction of CO, the evolution of CO_2 was promptly observed until the O_2^- species was completely consumed. This led the authors to conclude that the O_2^- species itself is the active species or O_2^- is a precursor to the active species. However, in Haruta's examination of Au/TiO₂ catalysts with EPR, it was possible to generate O_2^{-1} on titania in the absence of gold, which seems to speak against a mechanism requiring the presence of O_2^- (499). Furthermore, the authors note that after the catalyst has been exposed to both O₂ and CO, it can no longer stabilize O_2^{-1} species on the surface, although it can be argued that the anionic oxygen is quickly reacted with CO such that its observation is precluded.

Stable carbonate intermediates have been observed on the surface in several IR studies of real catalysts (191,358,375,377,403,421,422,430,473). Originally thought to be a key part of the mechanism (191,473), the prevailing view today is that CO_3 is a spectator species in the reaction (3,4,496) and that it can even act to deactivate the catalyst by blocking reaction sites (382,467,500-503). However, the transition could very well be a non-symmetrical carbonate such that preferential cleavage of the O-OCO bond is conceivable as has been proposed by both theorists and experimentalists (226,247,474).

Interestingly, although it has been hypothesized that the adsorption of oxygen, either dissociatively or molecularly, is the rate limiting step due to the extreme difficulty with which gold adsorbs oxygen, data from real catalysts seem to point in other directions. Haruta and others have consistently observed higher reaction orders for oxygen as for carbon monoxide. However, these reaction orders are frequently quite close to zero which would imply that both species are adsorbed to saturation on the surface at all times during the reaction and that the surface reaction itself is actually the rate limiting step (4,191). Bethke and Kung found that gold catalysts with larger particles were not only less active than those with smaller particles but also less selective with respect to CO oxidation when using a CO/ H_2 mixture (504). Since H_2 oxidation became favored for larger particles, one could conclude that the rate limiting step is not the dissociation of oxygen but rather the competition between hydrogen and CO adsorption that controlled the reaction rate.

Many studies of real catalysts have observed that the presence of water may enhance catalyst performance. Boccuzzi and Chorino suggested that water activates the dissociation of oxygen on gold particles and proposed that oxygen reacts with water to produce an oxygen radical available for reaction and two surface hydroxyl groups (505). Bond and Thompson (6), citing the difficulty of oxygen dissociation on gold surfaces, have even proposed reaction mechanisms involving water such as the one given below (and shown in Figure 29).

CO _{ad} + OH _{ad} → COOH _{ad}

$$O_{2 ad}$$
 + COOH $_{ad}$ \rightarrow $O_{2}H ad$ + CO_{2 g}

 $O_2H_{ad} + CO_{ad} \rightarrow OH_{ad} + CO_{2g}$

In this mechanism the requirement of breaking an O_2 bond in adsorbed oxygen is no longer necessary, as the O-OH bond is cleaved instead. The reaction is conceived via a reaction between a hydroxyl group that has moved from the support to a Au³⁺ ion and CO adsorbed on the gold particle. The resulting carboxylate group reacts with O_2^- and forms CO_2 and HO_2^- . The hydroperoxide can then react with a second carboxylate group which forms CO_2 and two surface hydroxyls, completing the cycle.

While some results have indicated that water acts to deactivate gold catalysts (359,403,432,506), the prevailing view is that water, at least in small amounts, substantially improves activity (3,4,487,507). In fact, in an ultra-dry environment, CO oxidation over Au/Mg(OH)₂ has been shown by Cunningham et al. to exhibit a negative activation energy (508). This led the authors to conclude that a direct reaction

Figure 29

Schematic representation of the early stage of the CO oxidation at the periphery of a gold particle involving the presence of hydroxyl groups and both Au^{3*} and Au^0 . See text for details. (6)

between hydroxyl groups on the surface and CO was a plausible reaction pathway. The presence of gold on the surface has been found to enhance the formation of hydroxyl groups on Au/Fe₂O₃ as compared to the bare support (366).

Haruta and Date have reported that in CO oxidation over Au/TiO₂, a maximum in activity occurred at 200 ppm H₂O (509,510) as shown in Figure 30. At high (6000 ppm) or low (0.1 ppm) concentrations of water the reaction rate was found to be a factor of 10 lower at 270 K. The improvement in CO conversion is not merely due to a concurrent water gas shift reaction because as pointed out by Sakurai *et al* (511) the water gas shift reaction for Au/TiO₂ is not active at room temperature.

Kung and co-workers have also found the presence of water to be beneficial to catalyst durability (512,513). Furthermore they have shown that Au/Al₂O₃ catalysts can be regenerated by either water or hydrogen with exposure to the former being more effective (514). They propose a mechanism similar to Bond and Thompson (6) whereby a CO is inserted in to a Au-OH bond, forming a hydroxycarbonyl on the surface. The hydroxycarbonyl can then react with oxygen as in the mechanism described above or alternatively, decarboxylation to CO₂ and Au-H can occur, followed by oxidation of the latter to regenerate the surface hydroxyl. Deactivation is believed to result from formation of carbonate groups which can be removed by reaction with water or hydrogen. Like Bond and Thompson, Kung and co-workers proposed that the presence of cationic gold is necessary for the reaction as cationic gold acts as the adsorption site for hydroxyls. Kung's group has also emphasized that chlorine contamination which displaces hydroxyl groups is a key factor in deactivation of gold catalysts (515).

Another possibility is that the coadsorption of water simply lowers the reaction barrier to CO oxidation. However, as Gong and Hu have recently pointed out in their examination of the catalytic role of H_2O on Pt(111), it is not likely that to be a primary pathway for CO oxidation as the reaction barrier (at least for Pt(111)) is on the same order as the barrier to water desorption (516).

With regard to the debate over the presence of metallic gold or cationic gold as the active species, Bond and Thompson proposed that both Au³⁺ and Au⁰ are required (6) as depicted in the schematic of Figure 29. In agreement with this type of mechanism, Gates and co-workers found that in active CO oxidation gold catalysts on MgO supports, both cationic and neutral gold species were present (517). Other researchers have since lent their support to mechanisms involving the formation of surface formate and the presence of both metallic and oxidic gold (450, 456). Bond and Thompson have further speculated that the mechanism may not be the same on all supports (6). Clearly, this is an area that demands more attention and should be an area where contributions from surface science can play a role.

It should be pointed out that CO oxidation is one of simplest reactions catalyzed by gold and that the understanding of other reactions such as the selective oxidation of propene to propylene oxide is even further behind. The reaction is optimized at about 100°C and requires the presence of hydrogen without which, the primary products are CO and CO₂ (10,518,519). Higher temperatures lead to the formation of secondary products such as propionaldehyde (520) and CO₂ while lower temperatures give very high selectivity although at the expense of activity (518). Particle size appears to be a key

Figure 30

Effect of the moisture concentration in the reaction gas on the CO oxidation reaction rate at 270 K over 1 wt.% Au/TiO₂ catalyst. (3 (data from 509))

Figure 31

The structure of the Au/titania interface for (a) anatase and (b) rutile demonstrating differences in the spacing between Au and Ti cations. (522)

variable as particles smaller than 2 nm favor the hydrogenation reaction to propane and large particles (521). The choice of support is of critical importance for selective oxidation of propene as only anatase TiO_2 supports and neither rutile nor amorphous TiO_2 supports have shown to be active (522). TiO_2 - SiO_2 mesoporous crystalline structures have proven to be active for this reaction (425,519,520,523,524) as well when possessing the proper Ti:Si ratio (generally below 3:100) (524,525). It is speculated that this may be related to a steric effect whereby Ti cations must be spaced at critical distances (522,523) as catalysts with such mesoporous material supports tend to be less active with increasing Au and Ti content (525). In Fig. 31, the relationship between Au and Ti cations for the rutile and anatase forms of titania are depicted, revealing the differences in spacing between Ti and Au (522). As discussed in Molina and Hammer's work with CO oxidation (474), the transition state is probably rather complex therefore very precise interatomic distances are required (523). As in the case of CO oxidation, some debate exists as to the nature of the active gold species (526). The mechanism is conceived to occur through the presence of a hydrogen peroxide intermediate which acts as the oxidizing agent (527). Theoretical calculations and experimental results alike have shown that formation of H₂O₂ is relatively facile over gold surfaces (528-530), lending credence to a mechanism involving hydrogen peroxide. Recent work has also confirmed the presence of a hydroperoxy species on the surface during a H_2 + O_2 reaction over Au/TiO₂ and it is thought that a similar species may form during propylene epoxidation (531). The use of D₂ induced an isotope effect, suggesting that the adsorption (or reaction) of hydrogen is the rate limiting step (518). In a recent review, Haruta speculated that after adsorption of oxygen on gold, the particles become electron deficient due to the electronegativity of oxygen and thereby adopt an electronic structure similar to platinum which allows for the facile dissociation of H_2 (532). However, the direct hydrogenation of propene to propane, which is observed for particles under 2 nm, is still difficult to explain in the context of such a mechanism.

In addition, very few studies have been performed that address the long-term performance of these catalysts (501,503,533-535). Some results have indicated that agglomeration of gold particles could pose a serious threat to long-term stable activity (192,394,412,421,476,535). Others have attributed the presence (403,432,506) or absence (507,508,512,513) of water as a contributing factor to deactivation. Minico et al ascribed deactivation of Au/Fe₂O₃ to an irreversible reduction of the gold from the 1+ state to metallic gold (378). For the selective oxidation of propylene, Au/TiO₂ catalysts also may experience problems with coking (527) as a result of propoxy decomposition at the acid sites of the titanosilicate support (523). Recently, it has been proposed that alkali metals such as lithium can be used as promoters to guard against sintering (536). While many researchers have reported phenomenal activity or selectivity these catalysts will not enjoy commercial success until they prove to be more robust.

6.3 Concluding remarks

Taken from a surface science perspective, real catalyst systems often seem to be poorly characterized and "standard" preparation techniques including the calcinations conditions vary widely from one lab to another leading to significant confusion in the analysis of results. Furthermore, experiments sometimes appear to be taken in an unsystematic manner and general trends are therefore often obscured. As one can readily appreciate from the preceding discussion in Section 6, in many cases the various results seem to be directly contradictory and therefore the reaction mechanism and active site for catalysis by gold nanoparticles remain unknown. It is also well known (but not often discussed) that the reproducibility of results of gold catalysts is of particular difficulty (426). Again, this largely stems from the lack of standardized catalyst preparation methods and the lack of proper characterization. Recently, certain standard catalyst preparations have been offered by Süd Chemie Japan in an effort to make comparison of results between various laboratories more meaningful (537). In the opinion of the current authors, surface science studies of well characterized model systems can make strong contributions in this regard as well. The approach we endorse involves the use of model systems whereby the addition of complexity in a stepwise manner allows one to easily identify the effects behind any changes in the chemistry observed. One criticism of surface science experiments is that the conditions under which these experiments are performed, i.e. ultrahigh vacuum, in no way replicate those conditions that the real catalyst must endure. In recent years, so called "high pressure" (1 mbar-1 bar) techniques such as SFG and in situ STM have been utilized in an effort (235, 538) to close the well-known pressure gap. Clearly, more experiments are required on model gold catalyst systems in reactive environments in order to better understand the complex chemistry of gold nanoparticles.

As we have previously discussed, a debate exists as to the nature of the active site and the mechanism. In general, CO is thought to adsorb on the gold particles themselves, probably on edge or step sites (3,4). The adsorption site of oxygen is less clear, although mechanisms involving oxygen adsorption on the support or at the particle-support interface are frequently invoked. Reaction is often conceived to occur at the interface and the nature of the reactive oxygen is still a mystery. In our opinion, molecular oxygen is likely to be the reactive species as theoretical calculations (197,216,247,474) consistently predict very high barriers to dissociation of oxygen, even for highly uncoordinated systems. Mechanisms involving hydroxyl groups largely stem from empirical evidence describing the improvement of catalytic performance in the presence of traces of water (507,509). Similar claims for a mechanism involving hydroxyls have been made previously for CO oxidation on alumina supported Pd (539). However, we have demonstrated in our lab that CO₂ does not form by CO exposure to Pd particles deposited on either pristine or hydroxylated alumina films (540).

Therefore, we would support a mechanism in which

molecular oxygen is adsorbed on highly uncoordinated sites on gold and reacts with CO, which is also adsorbed on gold. Recent work in our lab on the Au/FeO system indicates that quantum size effects alone do not control the adsorption of CO on gold nanoparticles and therefore suggest that the thickness of gold particles is not the critical parameter in CO oxidation (370). The requirement of a particular size therefore seems associated with the presence of highly uncoordinated atoms as has been demonstrated by Bahn et al (222). The increase of reactivity with decreasing coordination is not in itself surprising. However, the large differences in the chemistry of gold as its coordination number is reduced is exceptional and appears related to relativistic effects which do not arise in many other cases (34). The presence or absence of oxidic or anionic gold does not appear to be an important factor in CO oxidation, although there is evidence that Au^{+} could be critical for other reactions. As both theory and experiment have shown a strong preference for nucleation of gold on defects on oxide surfaces, the defect structure of the support is, therefore, critical in stabilization of the geometry required for the reaction. While it seems clear with the use of supports such as CeO₂ and Fe₂O₃ that a Mars-van Krevelen mechanism does come into play under certain reaction conditions, this is not the predominant mechanism. In conclusion, we believe that surface science studies of well-defined model systems can make strong contributions to the understanding of the unique reactivity of nanosized gold. It is our hope and intention to encourage other surface scientists to take on the challenges posed by this fascinating catalysis.

Acknowledgements

The authors would like to gratefully acknowledge Erika Popovich and Matthias Nachitzki for technical support. Furthermore, we would like to thank Prof. Claude Henry for his careful and thoughtful reading of the manuscript. We also are indebted to the Training Mobility and Research Project "Reactivity of clean and modified oxide surfaces" headed by Prof. Henry (TMR Marseille) and Training Mobility and Research Project "Catalysis by Gold" headed by Prof. Graham Hutchings (TMR AURICAT) for financial support as well as scientific discussions.

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