Monometallic Supported Gold Catalysts in Organic Transformations: Ring Making and Ring Breaking

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Abstract: Supported gold catalysts are highly active in oxidation reactions. Beside the most frequently studied CO oxidation, they are readily applied in the epoxidation of more or less complex olefinic compounds using air or oxygen directly or other oxidants like peroxides of various kinds. Less frequently though, the reverse reaction, ring opening with single or double C–O scission is also investigated. These and other ring making and breaking reactions are reviewed, and the catalytic roles of gold species are described.

Keywords: supported gold catalysts; oxygen-containing rings; ring synthesis; ring transformations

1. Introduction

Aerobic partial oxidations of alkanes and alkenes are important processes of the petrochemical industry. Selective oxidation is of huge importance in the synthesis of chemical intermediates and the epoxidation of alkenes by the electrophilic addition of oxygen to a carbon-carbon double bond remains one of the most significant challenges in oxidation catalysis. Although molecular oxygen is the most environmentally benign oxidant, in many cases far more reactive forms of oxygen are required to achieve reaction, and this can lead to by-products with a heavy environmental burden with respect to their disposal.
Following the success of ethylene epoxidation by silver-based catalysts [1–3], direct gas-phase propylene epoxidation using molecular oxygen has also been extensively studied. The propylene oxide (PO) selectivities were found to be very poor for silver due to the higher reactivity of propylene compared to ethylene. However, they are significantly better over gold-based catalysts; thus, it has been a promising research direction in recent years.

The topic of this review is this area widened in two respects. One is to include the epoxidation of other alkenes beside propylene and the other is to discuss the reverse reaction, the gold-catalyzed cleavage of the epoxide ring that is. We have done a literature search looking for aziridine or thiirane and higher N- or S-containing ring making and ring breaking reactions over supported gold catalysts, but papers like these could not be located. Neither could we find heterogeneous gold-catalyzed ring expansion or contraction reactions. These reactions either do not proceed or just have not reached the interest of catalytic chemists yet. Thus, the remaining area, i.e., epoxide ring making and breaking by heterogeneous gold-based catalysts, is the main topic of this review, but some ring making and ring transformation reactions other than the epoxide ring are also included. In every case, however, we restrict ourselves to monometallic gold, but attempting to cover all the support types used.

2. The Preparation of the Epoxide Ring

2.1. Epoxidation of Ethylene

The catalytic activities of Au (and also Ag and Au-Ag) deposited on various supports like on alumina, titania and ceria were studied in ethylene oxidation reactions [4] (Scheme 1).

Scheme 1. Epoxidation of ethane.

\[ \text{O}_2 \rightarrow \text{O} \]

The Au/TiO$_2$ catalysts proved to be the most selective towards ethylene oxide formation at relatively low ethylene conversion, and the Au/CeO$_2$ catalysts favored total oxidation over epoxidation even at very low temperatures. Beside the type of support the catalytic activities depended on the method of catalyst preparation governing the Au particle size and the interaction between the Au particles and the support.

2.2. Epoxidation of Propylene

Propylene oxide (PO) is an important chemical intermediate for the production of a variety of chemicals and polymers. Commercially, PO is produced via the chlorohydrin process and several hydroperoxide processes. However, the chlorohydrin process results in large amounts of chlorinated compounds, and the hydroperoxide processes typically generate stoichiometric quantities of co-products. The discovery by Haruta and co-workers [5] that nanoscale gold particles on titania supports provide a highly selective (\(\sim 99\%\)) route to vapor-phase PO production, with the use of a mixture of propylene, oxygen, and hydrogen under ambient pressure, has been a significant
breakthrough in heterogeneous catalysis. A direct, single-step epoxidation of propylene to PO has both practical and fundamental implications (Scheme 2).

**Scheme 2.** Epoxidation of propene.

Since then, among sporadic works of various researchers, five groups displayed intense activity. Haruta’s group remained a major player in this area [6–10]. In the epoxidation reaction they have used Au/Ti-MCM-41 and Au/TiO₂-SiO₂ catalysts with various Ti/Si molar ratios (0.25/100–6.0/100) [6]. Two different structure-directing organic templates (dodecyltrimethylammonium hydroxide and cetyltrimethylammonium hydroxide) were used in the synthesis of the Ti-MCM-41 support. TiO₂-SiO₂ mixed oxides were prepared by a sol-gel process. Gold nanoparticles were loaded by deposition-precipitation (DP) using various precipitating agents like LiOH, NaOH, KOH, RbOH, and CsOH. Au/Ti-MCM-41 with a Ti/Si = 3/100 molar ratio provided with the optimum performance in terms of conversion and PO selectivity: 3.2% and 93.5%, respectively at 373 K. At higher than 393 K PO selectivity decreased with an increase in the Ti/Si ratio. At 423 and 473 K, effective catalysts in terms of the above factors, had to have Ti/Si ratios of 2/100 and 1.5/100, respectively. A critical balance among Ti-content, Au loading and the reaction temperature was important to achieve higher propylene conversion, PO selectivity above 90% and lower H₂ (and O₂) consumption. Gold supported on TiO₂-SiO₂ mixed oxides showed very poor performance, however, PO selectivity appreciably increased with the increase in the calcination temperature of the support. IR and HRTEM analyses revealed that with the increase in the calcination temperature more isolated TiO₄ units with tetrahedral coordination was formed on the surfaces of Ti-SiO₂ supports. These isolated TiO₄ sites incorporated in the silica surface layers seemed to be an important structural factor for the selective deposition of Au particles and the selective PO formation [7].

After similar treatments a better performance was observed with Au/Ti-MCM-48 (initial conversion = 5.6%, PO selectivity = 92%), due to its three-dimensional pore system at 423 K [8]. GC-MS investigation of the extracted species by organic solvent from the used catalyst revealed that acidic as well as oligomeric species accumulated on the catalyst surfaces. These species were assumed to cause catalyst deactivation. Silylation of the catalyst Au/Ti-MCM-48 slowed down deactivation and helped to improve PO selectivity and decreased H₂ consumption as well.

Au/Ti-doped nonporous silica catalyst dried under vacuum at room temperature and calcined at 573 K in air, exhibited catalytic activity to form PO even at a temperature as low as 323 K [9]. The calcined catalysts were more stable and more active at higher temperatures than the dried catalysts. Simple filtration after aging the suspension resulted in catalysts more active and selective but less stable than the complete washing of the solid precursors. Pretreatment in argon increased the catalytic activity and H₂ efficiency appreciably over the Au catalysts deposited on Ti-doped nonporous silica as opposed to Au/Ti-MCM catalysts. Similar deactivation rate of Au catalysts supported on a variety of titania-silica supports with different porosities suggested that quick deactivation of Au catalysts within
hours was mainly affected by the surface properties rather than the pore structure and diffusion limitation of the supports.

It has been observed that in a H₂-O₂ mixture, the reaction of propene could be switched between hydrogenation and epoxidation over Au/Ti-based oxides [10] (Figure 1). Reaction pathways were strongly dependent on the size of Au particles and on the presence of alkalis. Hydrogenation prevailed over Au clusters smaller than 2.0 nm (in the absence of alkalis) and over Au nanoparticles larger than 5.0 nm. Alkali contamination could switch hydrogenation to epoxidation over Au clusters and promoted epoxidation over Au nanoparticles with diameters of 2.0–5.0 nm. It was also confirmed that the hydrogenation of propene was enhanced by the addition of a small amount of O₂.

**Figure 1.** The role of factors in deciding between the hydrogenation and epoxidation of propene. Reproduced with permission from Reference [10]. Copyright (2011) Elsevier.

Oyama *et al.* studied the oxidation of propylene to PO with H₂-O₂ mixtures over gold supported on mesoporous titanium silicates, Ti-TUD [11–13] and TS-1 [14]. The gold catalyst supported on Ti-TUD gave stable activity at low conversions of propylene (<6%) and high selectivity to PO (>95%) [11]. On the basis of kinetic data it was stated that the catalyst operated by the commonly accepted mechanism of hydrogen peroxide production on gold sites supported by *in situ* measured UV spectra, and epoxidation on titanium centers (Scheme 3). Carbon dioxide was formed primarily from further oxidation of PO rather than the oxidation of propylene, while water was produced from the reaction of hydrogen and oxygen. FTIR of the spent catalyst showed the formation of carbon species (probably a bidentate propoxy) due to propylene oxide decomposition, which is a probable cause for catalyst deactivation. Surface formate and acetate species were also observed as oxidation products of the bidentate species in the spent catalyst.

7% Au loading could be achieved on promoting the Ti-TUD support with Ba²⁺ at pH 7. The Au particle size was 2.0 nm by transmission electron microscopy, whereas at pH 9 much lower Au loading of 0.11 wt.% was produced with particle size of about 0.9 nm estimated by X-ray absorption fine structure measurements. At 423 K and 0.1 MPa total pressure, the catalyst prepared at pH 7 gave a steady-state propylene conversion of 2.1%, a propylene oxide selectivity of 79%, and a H₂ efficiency of 3.8%, whereas that prepared at pH 9 gave a conversion of 1.4%, a propylene oxide selectivity of 99%, and a H₂ efficiency of 17%. It is concluded that very small Au particles (about 1 nm) were the
most active for epoxidation, whereas larger Au particles (about 2 nm) were less active because they promote direct \( \text{H}_2 \) oxidation to \( \text{H}_2\text{O} \). X-ray absorption near-edge spectroscopy results indicated that under reaction conditions, the small particles had partially oxidized gold but the larger particles had metallic gold, suggesting that the smaller particles had high coverage of oxygen or oxygen-derived species [12].


A series of Au/titanium silicalite-1 (TS-1) catalysts with different Si/Ti ratios and promoted with alkali and alkaline earth cations were prepared by the DP technique and tested for direct propylene epoxidation. It was found that the gold loading and catalytic activity were highly dependent on the \( \text{pH} \) of the DP synthesis solution and the final composition of the catalyst. Addition of the cations of group 1 metals such as K or Cs had little effect on the gold content, but increased the activity, while those of group 2 metals such as Mg, Ca, Sr, and Ba increased both the gold content and the catalytic activity. The highest improvement was provided by a Mg\(^{2+}\)-promoted catalyst giving 50% enhancement of activity at 443 K and 0.1 MPa with a \( \text{H}_2/\text{O}_2/\text{C}_3\text{H}_6/\text{Ar} = 1/1/1/7 \) feed mixture. Ammonia temperature-programmed desorption (NH\(_3\)-TPD) measurements indicated little change in adsorption amount with promotion indicating that the yield increase was not due to the elimination of acidic sites on the catalyst. Instead, the improved catalytic performance was ascribed to increased Au capture efficiency and dispersion by the catalyst [13].

Cyanide treatment of Au/TS-1 provided with unexpected results. Catalysts treated with dilute solutions of sodium cyanide resulted in preferential removal of small gold particles, while catalysts treated with concentrated solutions resulted in dissolution of gold and its re-precipitation as Au(I) cyanide. X-ray absorption spectroscopy demonstrated that catalysts that produce PO in the presence of hydrogen and oxygen mixtures had supported Au(III) oxide nanoparticles of 3 nm size after synthesis, which were reduced to gold metal under reaction conditions. Samples treated with concentrated solutions of sodium cyanide resulted in supported Au(I) cyanide particles of large size (9–11 nm). These particles did not produce PO, but, surprisingly, showed high selectivity toward propylene hydrogenation. Increasing Au(I) cyanide particle size resulted in a decrease in hydrogenation activity [14].

Propane epoxidation has also been attempted (Figure 2).
It was carried out by sequential propane dehydrogenation, propylene epoxidation steps using a two-catalyst bed and H\(_2\) and O\(_2\) as the oxidant mixture. The propane dehydrogenation step used an Au/TiO\(_2\) catalyst that was active at low temperature (443 K), while for the propylene epoxidation step an Au/TS-1 catalyst was applied. *In situ* Au L3-edge X-ray absorption near-edge (NEXAFS) structure and UV-visible measurements on Au/TiO\(_2\) under propane dehydrogenation conditions showed activation of oxygen on gold nanoparticles and evidence for the formation of adsorbed oxygen intermediate species responsible for the production of propylene. Propane epoxidation with H\(_2\) and O\(_2\) at 443 K and 0.1 MPa with the dual Au/TiO\(_2\) and Au/TS-1 catalysts resulted in an overall propane conversion of 2%, propylene selectivity of 57%, and PO selectivity of 8%. The catalysts showed little deactivation and maintained their conversion and selectivity levels for the 12 h duration of the measurements [15].

Delgass and co-workers have also been deeply involved in studying propylene epoxidation over gold-based catalysts [16–20]. The epoxidation reaction was investigated over gold particles prepared by the DP method on various modified titanium silicalite-1 (TS-1) supports over a reaction time of 24–36 h in a flow reactor at temperatures of 413, 443, and 473 K. Gold deposition at pH 9–10 allowed for a consistent amount of 1–3 wt.% of the gold available in solution to be deposited, while still maintaining gold particle diameters in the 2–5 nm range, as observed by TEM [16]. These Au/TS-1 catalysts achieved propylene conversions of 2.5–6.5% and PO selectivities of 60–85% at 443 K. A key result of the work is that PO rates were not highly influenced by the TS-1 particle size and were thus not proportional to the specific external surface area of the support. The conclusion that activity may have resided in the channels of the TS-1 was supported by the finding that the observable gold particles decorating the TS-1 particles only accounted for ca. 30% of the total gold content of the catalyst. Increasing the gold loading up to 0.74 wt.% did not increase PO rates proportionally, suggesting that the active Au-Ti PO forming centers were limited. In contrast to the prevailing interpretation of this catalyst that a critical Au
particle diameter of 2–5 nm is essential for PO activity, the results were consistent with a molecular cluster model where extremely small gold clusters are located near Ti sites inside the TS-1 pores or on the external surface. These clusters were thought to be active for propylene epoxidation. Treatment of the calcined TS-1 support with 1 M NH₄NO₃ at 80 °C, followed by vacuum drying, produced a modified TS-1 support material having a four-fold increase in Au capture efficiency and produced catalysts with 5–10% conversion of propylene with 75–85% selectivity for PO at 200 °C in a flow reactor [17].

The superior activity and stability of Au/TS-1 catalysts allowed the first comprehensive kinetic analysis of the propylene epoxidation system in the absence of significant deactivation. A unique design of experiments combining the best features of factorial experiments with one-at-a-time experimentation over the nonflammable range was used to collect kinetic information, from which a power rate law was extracted. Explaining the resultant fractional reactant orders \(O_2 = 0.31 \pm 0.04\), \(H_2 = 0.60 \pm 0.03\), and \(C_3H_6 = 0.18 \pm 0.04\) required a sequence of elementary kinetic steps having a minimum of two active sites participating in the rate-determining step [18,19]. A reaction sequence was proposed that accounted for the experimentally determined reaction orders and was consistent with DFT calculations [20] (Figure 3). This mechanism suggests that titanium and gold sites must generate and use the epoxidation oxidant simultaneously rather than sequentially, as previously suggested in the literature (Figure 3).

**Figure 3.** The transition state geometry for attack of H₂O₂ on Au₃/T6-Ti-defect site to form Ti-OOH species inside the TS-1 pores. All of the MM atoms and some atoms in the QM region were removed for clarity. The atomic distances in Å are indicated. Reproduced with permission from Reference [20]. Copyright (2007) American Chemical Society.

The Weckhuysen group synthesized highly dispersed gold nanoparticles within the channels of a mesoporous Ti-SBA-15 support, followed by thorough catalyst characterization and testing in the selective epoxidation of propene to PO [21]. For this purpose, two series of Ti-SBA-15 materials differing in their Ti content were prepared by either grafting or direct synthesis. It was found that the Au/Ti-SBA-15 materials obtained by Ti grafting had higher catalytic activity than the samples in
which Ti-SBA-15 was obtained by direct synthesis. These differences in catalytic behavior were attributed to differences in the amount and dispersion of Ti within the mesoporous silica support as well as to differences in the size of Au nanoparticles. The adsorption of propene on supported gold nanoparticles has been experimentally identified as a reaction step in the hydro-epoxidation of propene [22]. This new finding was made possible by applying a detailed analysis of in situ measured XANES spectra. For this purpose, Au/SiO₂ catalysts were investigated since this support was more inert and propene was not converted. Propene adsorption was investigated by using the oxidation of hydrogen as probe reaction. It was shown that co-feeding of propene dramatically decreased the hydrogen oxidation rate. Since it has been reported in the literature that the oxidation of hydrogen occurs exclusively over gold nanoparticles, this inhibition by propene can be attributed to adsorption of propene on the gold nanoparticles. Analysis of the in situ XANES spectra confirmed the adsorption of propene on the gold nanoparticles and the mode of adsorption was determined to be π-bonding. Comparative experiments with ethene and propane confirmed this π-bonded adsorption, since ethene similarly inhibited the oxidation of hydrogen, while propane had only a minor effect.

Titanium-containing hexagonal mesoporous silicas (Ti-HMS) with wormhole structure and Si/Ti molar ratios ranging from 10 to 40 have also been studied [23]. Catalytic results showed that the Au/Ti-HMS catalyst exhibited superior performance in terms of propylene conversion, PO selectivity, and H₂ efficiency in comparison with the Au catalysts supported on the conventional Ti-containing mesoporous materials. Beside the Si/Ti molar ratio, the chain length of alkylamine for the Ti-HMS preparation was crucial for the enhancement of catalytic performance. Specifically, 9.0% of propylene conversion, 97.3% of propylene oxide selectivity, and 30.4% of H₂ efficiency could be obtained at 373 K in the initial 30 min of time-on-stream on the Au/Ti-HMS catalyst. The Ti-HMS having a Si/Ti molar ratio at 20 was prepared by using tetradecylamine as the templating agent. Regeneration of the spent catalyst by calcination in air gave almost no change in PO selectivity but about 25% loss in propylene conversion. The enhanced catalytic performance of Au/Ti-HMS catalyst may be essentially attributed to the homogeneous dispersion and uniformity of titanium species in combination with accessible pore structure.

The works of all groups indicate that in the preparation of gold catalysts that are highly selective in PO formation Ti-containing supports must be used, the particle size of Au is decisive, to achieve its optimum the deposition-precipitation technique is the best and the alkali content of the catalyst also influences catalytic activity. The calcination temperature is important as well and the reaction temperature should not be too high.

2.3. Epoxidation of Styrene

The epoxidation of styrene proceeds in the liquid phase over supported gold catalysts using peroxides, mostly tert-butyl hydroperoxide, as co-reactant (Scheme 4).

**Scheme 4.** Epoxidation of styrene.
In Choudry’s group many different oxides (MgO, CaO, SrO, BaO, Al₂O₃, Ga₂O₃, In₂O₃, Tl₂O₃, TiO₂, Cr₂O₃, MnO₂, Fe₂O₃, CoO, NiO, CuO, ZnO, Y₂O₃, ZrO₂, La₂O₃, Ce₂O₃, Nd₂O₃, Sm₂O₃, Eu₂O₃, Tb₂O₃, Er₂O₃, Yb₂O₃ and U₃O₈) as support have been tested [24–28]. The supported catalysts were prepared by depositing gold on the support by the DP and the homogeneous DP methods. The catalysts were calcined at various temperatures (400 °C–900 °C) and the catalytic activity of the supported nanogold catalysts in the epoxidation of styrene to styrene oxide by tert-butyl hydroperoxide were correlated with the Au loading and/or Au particle size of the catalysts. The reaction was strongly influenced by a number of parameters, such as the metal oxide support, the method of gold deposition, the gold loading as well as the catalyst calcination temperature. For characterizing the catalytic performance (both the activities and the selectivities) some activity orders (the selectivity values changed in the same sequence) could be set up, like Au/MgO > Au/Tl₂O₃ > Au/Yb₂O₃ > Au/Tb₂O₃ > Au/CaO (or TiO₂); Au/Tl₂O₃ > Au/In₂O₃ > Au/Ga₂O₃ > Au/Al₂O₃. The nanogold particles-support interactions seem to play an important role in controlling the deposition of gold (amount of gold deposited and size and morphology of gold particles), formation of different surface gold species (Au⁰, Au⁺ and Au³⁺), consequently, they control the catalytic performance (both the activity and selectivity) of the supported nanogold catalysts [25–28].

Three kinds of mesoporous alumina (denoted as meso-Al₂O₃-x (x = a, b, c)) supports with different surface basicities were synthesized by various structure-directing agents and different assembly pathways [29]. It has been found that the mesoporous alumina had more abundant surface basic sites than γ-Al₂O₃. Au nanoparticles were deposited on these supports via the homogeneous DP method using urea as the precipitating agent. The dispersion and average size of the gold particles were found to be dependent on the number of surface basic sites on the supports. Transmission electron microscopy (TEM) observations showed a homogeneous distribution of gold particles lower than 4 nm on meso-Al₂O₃-b and meso-Al₂O₃-c having more basic sites on the surface. XPS spectra revealed that only metallic gold was present on the supports irrespective to the surface properties. The catalysts were employed as highly active/selective and reusable catalysts for the epoxidation of styrene with anhydrous tert-butyl hydroperoxide. Au nanoparticles as well as the surface basic sites of the support were thought to be responsible for the epoxidation.

Somewhat unusual supports were also used in preparing the gold-based catalysts like (S)-(−)-2-pyrrolidinone-5-carboxylic acid (Py)-modified SBA-15 [30], S-containing organic-inorganic hybrid mesoporous silicas [31] or carbon nanotubes [32].

Highly dispersed Au nanoparticles could be deposited in the mesopores of (S)-(−)-2-pyrroldinone-5-carboxylic acid (Py)-modified SBA-15 (Au/SBA-15-Py) [30] (Figure 4). ¹³C NMR and IR spectroscopies indicated that Py species were successfully grafted on the surface of mesopores in SBA-15; XRD patterns and N₂ adsorption isotherms showed that the mesostructures were well preserved; TEM images clearly confirmed the uniform Au nanoparticles in the mesopores; and XPS suggested an interaction between Au nanoparticles and Py species. Interestingly, and importantly, Au/SBA-15-Py catalysts always exhibited superior catalytic properties in the oxidation of cyclohexene and styrene by molecular oxygen at atmospheric pressure, compared with the pyrrolidone-free SBA-15 supported Au catalyst (Au/SBA-15-N). This phenomenon was reasonably related to the interaction between Au nanoparticles and Py species, which was consistent with results of density functional theory (DFT) calculations.

For the mesoporous organosilica it was found that the location and structure of organic moieties in the mesostructures played a crucial role in the dispersion level of gold nanoparticles, and thus, had a significant effect on the catalytic performance. Under optimum conditions, the gold nanoparticles supported on a periodic mesoporous organosilica with bridging disulfide-ionic liquid moieties (Scheme 5) showed excellent catalytic performance and reusability in the epoxidation of styrene in the presence of H$_2$O$_2$–acetonitrile [31].

Scheme 5. The bridging organosilica precursor.

\[
\text{[(EtO)$_3$Si]_2N}_3\text{((CH$_2$)$_3$} \quad \text{[(EtO)$_3$Si]_2N}_3\text{((CH$_2$)$_3$}
\]

The catalytic activity of carbon nanotube supported gold catalysts prepared by the DP method using urea for precipitation was examined for the oxidation of styrene using tert-butyl hydroperoxide as oxidant. The system showed good epoxide selectivity. The other factors, such as solvent, reaction time,
concentrations of oxidant and catalyst, have also been investigated and reaction conditions were optimized. It is a novel highly active/selective and reusable heterogeneous catalyst for styrene epoxidation [32].

2.4. Epoxidation of Stilbene

The stilbene epoxidation reaction can be efficiently catalyzed by supported gold nanoparticles (Scheme 6).


A free-radical mechanism has been evidenced in the liquid-phase stereoselective epoxidation of trans-stilbene using methylycyclohexane as solvent, limited amount of tert-butyl hydroperoxide, and supported gold catalysts [33,34]. Trans-stilbene oxide was the major reaction product observed, with selectivities up to 88% over the Au/TiO$_2$ reference catalyst. However, the selectivity decreased significantly on using Au/C instead of oxide-supported gold catalysts or H$_2$O$_2$ instead of tert-butyl hydroperoxide. It seemed that tert-butyl hydroperoxide was the radical source, while methylycyclohexane was propagating the active radical. XPS measurements showed the presence of Au$^0$ (90%) and Au$^+$ (10%) on the Au/C catalyst and Au$^{5-}$ (90%) and Au$^+$ (10%) on the Au/TiO$_2$ catalyst. Both gold and, to a minor extent, titania seemed to be involved in the reaction cycle.

Citrate-functionalized titania nanocrystallites were synthesized from a heteroleptic titanium alkoxide precursor in a low-temperature hydrolytic process and used as gold catalyst support for CO oxidation and aerobic stilbene epoxidation [35].

The support had little influence on the intrinsic activity of gold but more on the apparent reaction rates which are a combination of catalytic activity and diffusion limitations. These were here minimized by using gadolinium-doped titania nanocrystallites as support for gold nanoparticles [36]. This material was obtained by the mild hydrolysis of a novel Gd$_4$TiO(OiPr)$_4$ bimetallic oxo-alkoxide. It led to enhanced wettability of the <3 nm gold particles in the tert-butyl hydroperoxide initiated epoxidation of stilbene in methylycyclohexane. The rate-determining step of this reaction was identified as the gold-catalyzed homolytic decomposition of tert-butyl hydroperoxide generating radicals and initiating the methylycyclohexane-mediated epoxidation of stilbene, yielding a methylycyclohexan-1-ol/trans-stilbene oxide mixture.

The preparation and characterization and catalytic testing in the aerobic oxidation of cyclohexene and trans-stilbene in the liquid phase of highly dispersed gold nanoparticles in ordered mesoporous carbons have also been reported [37]. The carbon supports were prepared using gold-containing functionalized SBA-15 silica templates. Two series of Au/SBA-15 templates were made. In the first
series the size of the gold particles was determined by the pore size in the ammonium-functionalized silicas, while the mercaptopropyltrimethoxysilane grafting agent allowed the possibility of controlling the particle size inside the mesopores. Both series provided with highly ordered mesoporous carbons with gold particles incorporated in the carbon nanorods. Their catalytic activities and selectivities in epoxidation were appreciable and practically did not depend on the support.

2.5. Epoxidation of Cyclooctene

The solvent-free epoxidation of cis-cyclooctene with air using supported Au catalysts with small amounts of a hydroperoxide was performed [38,39] (Scheme 7). In the absence of a hydroperoxide initiator, using air at atmospheric pressure, no reaction was observed. Choice of the peroxide initiator proved to be crucial, and in the absence of a catalyst or a support, the reaction of the alkene could be observed with [(CH₃)₃CO]₂, cumene hydroperoxide and tert-butyl hydroperoxide only when high concentrations were used at high temperatures. At temperature of 80 °C tert-butyl hydroperoxide was found to be the most selective to epoxide formation. In contrast, cumene hydroperoxide was highly reactive under all conditions evaluated. tert-Butyl hydroperoxide was selected for more detailed study.

Scheme 7. Epoxidation of cis-cyclooctene.

Using mild solvent-free conditions the hydroperoxy initiator persisted in solution for only a few minutes, being initially adsorbed on the catalyst surface. Subsequently, it decomposed to establish a reactive species that could propagate the selective oxidation process. The observation of an induction period might in part be due to the adsorption of the radical initiator blocking surface sites as well as the establishment of the reactive species. Use of graphite as a support gave the best combination of selectivity and conversion. In general, the selectivity to the epoxide increased with reaction temperature from 60–80 °C and was the highest at 80 °C. Other carbon supports, e.g., activated carbon, were less effective. TiO₂- and SiO₂-supported Au catalysts were also selective for the epoxidation, and the general order of activity was: graphite > SiO₂ > TiO₂. The major by-product was allylic alcohol. The preparation of the catalysts by the sol-immobilization method enhanced catalyst activity with retention of selectivity to the epoxide [38]. Extensive studies were also reported concerning the reusability of the gold/graphite catalyst. The catalyst was found to be inhibited by the epoxide product but it was demonstrated that the effect of this was negligible for reused catalysts over a long reaction time [39].

3. Ring-Opening Reactions of Propylene Oxide

3.1. Single and Double C–O Scissions

An Au/MgO catalysts with 3% Au loading was found to be active in the ring-opening reaction of PO as low as 363 K, irrespective to the method of pretreatment, although the level of activity was largely dependent on pretreatment conditions [40] (Scheme 8).
The main reaction at 363 K and 393 K was deoxygenation (scission of both C–O bonds) producing propylene (2). At 423 K the sum of acetone (3) and 2-propanol (5) concentrations (the products of isomerization and hydrogenative ring opening, respectively) nearly reached that of propylene, and at 473 K the single C–O scission reaction became the major transformation pathway.

Desorbed water (7) appeared in most cases at 393 K, although in small quantities. Its concentration increased with the increase in temperature. As it has been indicated, the activity of the catalysts was dramatically influenced by the method of pretreatment. When it was finished with hydrogen the conversion values were always above 70%, but often they were in the 90+% range.

A 3 wt.% Au/multiwall carbon nanotube (MWNT) composite prepared by sonication-assisted deposition, followed by reduction in H₂ flow, was active in transforming PO in a H₂ atmosphere [41]. Deoxygenation was the major transformation pathway in the 363–473 K temperature range. Let us note, however, that isomerization (already at 363 K) and from 393 K hydrogenation (products of single C–O scission) also took place. Hydrogenation became important at higher temperatures. Similar behavior was observed over a 3 wt.% Au/activated carbon catalyst albeit the transformation started at 423 K and became appreciable only at 473 K. The 3 wt.% Au/graphite catalyst became active at a temperature as high as 473 K and double C–O scission only took place producing propylene [42].

Unsupported gold was only slightly active even at 473 K and deoxygenation was inferior to isomerization. The transformation spectrum was also much narrower. It was clear that the supports played important role not only by dispersing the metal, but also allowing traffic of certain species, probably oxygen from the metal. Possibly, the metal–support interface played crucial role in this reaction. Crystal defects might have been formed even on the seemingly perfect graphite sheet (Figure 5)—let alone the MgO support—during sonication, hydrogen reduction, and upon methylloxirane transformations.
3.2. Selective Deoxygenation: Double C–O Scission

Selective transformation of epoxides into the corresponding alkenes is an important reaction because it allows the use of oxirane rings as protecting groups for carbon–carbon double bonds. However, if this reaction is performed using stoichiometric amounts of reagents, it will produce large amounts of undesirable waste. The development of recyclable, appreciably active and very selective heterogeneous catalyst would be an environmentally benign and highly desirable alternative. Ideally, molecular hydrogen (H₂) should be used as a reducing reagent because, theoretically, water is the only by-product. Unfortunately, the use of H₂ often results in nonselective reduction of epoxides yielding alcohols and alkanes as by-products.

Recently, it was shown that hydrotalcite-supported [HT (hydrotalcite): Mg₆Al₂(OH)₁₆CO₃·nH₂O] gold and silver nanoparticle catalysts have high activities for the deoxygenation of many different epoxides to alkenes (Table 1) with >99% selectivity, using 2-propanol as an environmentally friendly reductant [43].

Furthermore, CO/H₂O was found to work as an alternative reductant for the selective deoxygenation of epoxides to alkenes in water under mild reaction conditions [44]. The same group also demonstrated that the Au/HT could act as a highly efficient heterogeneous catalyst for the deoxygenation of a large variety of epoxides to alkenes with H₂ used as an ideal reducing agent.

The selectivities for alkenes were over 99% at high conversions. After the reaction, solid Au/HT could be easily recovered from the reaction mixture and reused with no decrease in its catalytic efficiency [45]. The possible transformation mechanism is depicted in Figure 6.

Very recently, the preparation of Au/TiO₂ catalysts with very small gold nanoparticles (mean particle size: 1.9 nm) has been described, and were used for catalyzing the deoxygenation reaction even under sub-ambient temperatures mediated by CO and H₂O [46]. The CO-mediated deoxygenation system had the following advantages: (i) unprecedented high activity under very mild conditions; (ii) high selectivity and functional group tolerance; (iii) the use of safe and easy-to-handle catalysts and reducing reagents; and (iv) a simple workup procedure, namely catalyst/product separation by filtration.
Table 1. Deoxygenation reactions of various epoxides over Au/HT (or Ag/HT) catalyst \[^a\]. Reproduced with permission from Reference [43]. Copyright (2010) Wiley-VCH Verlag GmbH & Co. KGaA.

<table>
<thead>
<tr>
<th>Entry</th>
<th>Substrate</th>
<th>Product</th>
<th>Catalyst</th>
<th>t (h)</th>
<th>Yield (%) [^b]</th>
<th>Sel. (%) [^b]</th>
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<td>1</td>
<td></td>
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<td>Au/HT</td>
<td>4</td>
<td>99</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>2 [^c]</td>
<td></td>
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<td>4</td>
<td>99</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>3 [^d]</td>
<td></td>
<td></td>
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<td>4</td>
<td>97</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>4 [^e]</td>
<td></td>
<td></td>
<td>Au/HT</td>
<td>72</td>
<td>95</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>5</td>
<td></td>
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<td>Ag/HT</td>
<td>8</td>
<td>95</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>6 [^c]</td>
<td></td>
<td></td>
<td>Ag/HT</td>
<td>8</td>
<td>93</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>7 [^d]</td>
<td></td>
<td></td>
<td>Ag/HT</td>
<td>8</td>
<td>93</td>
<td>(&gt;99)</td>
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<tr>
<td>8</td>
<td></td>
<td></td>
<td>Au/HT</td>
<td>4</td>
<td>99</td>
<td>((E/Z = 2:3))</td>
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<tr>
<td>9</td>
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</tr>
<tr>
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<td>98</td>
<td>(&gt;99)</td>
</tr>
<tr>
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<td>8</td>
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<td>(&gt;99)</td>
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<td>6</td>
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<td>(&gt;99)</td>
</tr>
<tr>
<td>13 [^g]</td>
<td></td>
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<td>24</td>
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<td>(&gt;99)</td>
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<tr>
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<td>89</td>
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<tr>
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<td></td>
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<td>trace</td>
<td>(&gt;99)</td>
</tr>
<tr>
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<td></td>
<td>Au/HT</td>
<td>24</td>
<td>87</td>
<td>(&gt;99)</td>
</tr>
<tr>
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<td>Ag/HT</td>
<td>24</td>
<td>trace</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>25 [^f]</td>
<td></td>
<td></td>
<td>Au/HT</td>
<td>4</td>
<td>97</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>26 [^g]</td>
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<td>Au/HT</td>
<td>48</td>
<td>93</td>
<td>(&gt;99)</td>
</tr>
<tr>
<td>27 [^f]</td>
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<td></td>
<td>Ag/HT</td>
<td>24</td>
<td>trace</td>
<td>(&gt;99)</td>
</tr>
<tr>
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<td></td>
<td>Au/HT</td>
<td>12</td>
<td>81</td>
<td>((E/Z = 1:1))</td>
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<tr>
<td>29 [^b]</td>
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<td>30 [^b]</td>
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<td>(&gt;99)</td>
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<td>31 [^b]</td>
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<td></td>
<td>Ag/HT</td>
<td>24</td>
<td>trace</td>
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Table 1. Cont.

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<tr>
<th></th>
<th>Reaction conditions: Catalyst (0.1 g), substrate (1 mmol) toluene (5 mL), 2-propanol (0.6 mL);</th>
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<tbody>
<tr>
<td>32</td>
<td>Au/HT</td>
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<tr>
<td>33</td>
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<tr>
<td>37</td>
<td>Ag/HT</td>
<td>24</td>
<td>trace</td>
</tr>
</tbody>
</table>

[a] Reaction conditions: Catalyst (0.1 g), substrate (1 mmol) toluene (5 mL), 2-propanol (0.6 mL); [b] Determined by GC and LC methods using an internal standard technique; [c] Cycle 1; [d] Cycle 2; [e] Conditions: catalyst (0.02 g), substrate (20 mmol), 2-propanol (20 mL); [f] 80 °C; [g] Conditions: catalyst (0.02 g), substrate (20 mmol), 2-propanol (20 mL); [h] Substrate (0.3 mmol), catalyst (0.2 g).

Figure 6. Heterolytic dissociation of H\textsubscript{2} over basic sites (BS) and deoxygenation of the epoxide over the gold nanoparticles. Reproduced with permission from Reference [45]. Copyright (2011) Wiley-VCH Verlag GmbH \\& Co. KGaA.

4. Preparation and Transformation of Rings Other than the Epoxide Ring

Several heterogenized catalytic systems based on Au have been prepared and analyzed and their efficiencies were evaluated in the cycloisomerization of \(\gamma\)-acyltylenic carboxylic acids to valuable \(\gamma\)-alkylidene \(\gamma\)-butyrolactones (Scheme 9). Among the heterogeneous catalysts (Au/CeO\textsubscript{2}, Au/MgO, Au/TiO\textsubscript{2}, Au/\(\beta\)) prepared, the first two were not active at all, Au/TiO\textsubscript{2} displayed fair, and Au/\(\beta\) outstanding activities [47].

Scheme 9. Supported gold catalyzed transformation of a \(\gamma\)-acyltylenic carboxylic acid.

\(\alpha\)-Pinene was isomerized to camphene over a 2.2 wt.% Au/\(\gamma\)-Al\textsubscript{2}O\textsubscript{3} catalyst in the 463–483 K range using a solution of the reagent in n-octane as the initial reaction mixture and H\textsubscript{2} or N\textsubscript{2} as a carrier gas. This reaction representing alkyl group migration from one carbon to a neighboring carbon.
(Wagner-Meerwein rearrangement) has high practical significance as a first step in the industrial synthesis of camphor from \( \alpha \)-pinene (Scheme 10).

**Scheme 10.** The \( \alpha \)-pinene camphene transformation over Au/\( \gamma \)-Al\(_2\)O\(_3\) catalyst.

Under these conditions, the selectivity to camphene reaches 60–80% at 99.9% conversion of \( \alpha \)-pinene. Prominent catalyst deactivation has been observed at increased \( \alpha \)-pinene concentrations in the inlet reaction mixture due to carbonaceous deposits that blocked the catalyst surface. The catalysts could be regenerated completely applying \( \text{O}_2 \) flow at 923 K [48].

5. Conclusions and Outlook

Supported gold catalysts are active in ring making and breaking reactions when the ring contains oxygen. Gold deposited on various Ti-containing supports are excellent propylene epoxidation catalysts in the laboratory, and they are very promising as industrial catalysts as well. Ring opening as well as deoxygenation reactions are useful tools in the hands of the preparative chemists. The extensive studies revealed that for every kind of reaction, the supports were crucial parts of the catalysts; unsupported gold was hardly active or not active at all.

Current activities in this area are concentrated on making and breaking oxygen-containing rings, however, it would be nice to see if the preparation and transformation of ring systems containing heteroatom(s) other than oxygen were available through heterogeneous gold catalysis.

Acknowledgment

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References


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