Ceria-Based Mixed Oxide Supported Nano-Gold as an Efficient and Durable Heterogeneous Catalyst for Oxidative Dehydrogenation of Amines to Imines Using Molecular Oxygen

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Abstract

The present work is intended to determine the catalytic activity of Mixed Oxide supported gold for aerobic oxidative dehydrogenation of amines to imines using Ceria as a main constituent of the each support. The model catalysts Au/CeO2:TiO2 Au/CeO2:SiO2, Au/CeO2:ZrO2 and Au/CeO2:Al2O3 were prepared by deposition co-precipitation method and deposition of gold was determined by EDEX analysis. The supported nano-gold catalyzes the dehydrogenation of secondary amines to imines without loss of activity. On recycling good amount of product yield is obtained. Oxidation of secondary amines to imines is carried at 100˚C and almost 90 % conversion was obtained with >99% selectivity. © 2012 BCREC UNDIP. All rights reserved

Keywords: Heterogeneous Catalyst; Nano-Gold; Metal Oxides; Imines


1. Introduction

Imines and their derivatives have long been recognized as basic building blocks for the synthesis of nitrogen heterocycles, especially in the arena of alkaloid synthesis basic building blocks and various synthetic routes for the synthesis of imines are reported. Although the synthesis of aldimes is quite easy by using aldehyde and primary amines, the synthesis of many classes of imines is problematic e.g. ketimines (especially aryl ketimines) [1-3]. The synthesis of ketimines is difficult and requires metal chloride, which is used as dehydrating agent to overcome the competing reverse reaction, but metal chlorides generate HCl which leads to unwanted side reactions in many systems even in the presence of excess amine [4, 5]. Stoichiometric methods utilizing strong oxidants such as Swern oxidation have the disadvantage of forming major amounts of side products [6, 7]. So the oxidative dehydrogenation of secondary amines has been explored as an alternative route for the preparation of imines.

The discovery of the catalytic properties of gold nanoparticles is one of the most exciting contributions to catalysis and has led to the renaissance of gold chemistry. Gold based catalysts attracted wide attention ever since Haruta et al., found that supported gold catalysts are highly

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active for the oxidation of CO at temperatures as low as -73 °C when deposited as nanoparticles on metal oxides [8,9]. The catalytic activity of a gold catalyst does not depend on the size of the gold particles only, but also on the nature of the support material, the preparation method, and the activation procedure [10-13]. Supported gold nanoparticles have attracted considerable attention because of their extraordinarily high activity and selectivity. Support materials like transition metal oxides provide high reactive surface area, thermal stability, higher dispersion of nano-metallic particles and longer catalyst life etc [14]. The discovery that Au clusters dispersed on metal oxide supports act as efficient catalysts for a variety of low-temperature reactions has led to a great deal of study, with the hope that supported Au clusters may someday replace the less active catalysts currently in use. The use of gold as a catalyst requires careful and unconventional preparation methods for achieving a very small gold particle size. These nano-gold catalysts provide fast reaction rate, high selectivity, and low reaction temperature [15]. The exceptional catalytic capability of supported gold is associated with the size and shape of the nanoparticles, the degree of coordinative unsaturation of the gold atoms, and the interactions between gold and the oxide support [16].

CeO₂ is having many beneficial properties in catalysis, such as improving the dispersion of surface metals, store and release oxygen. The later property is expected to help minimizing the pyrophorocity of the metal supported on CeO₂ [17]. The two unique features which are responsible for making CeO₂ a talented substance for exercise either as a support or as an active catalyst are (a) the Ce³⁺/Ce⁴⁺ redox couple, with its ability to shift between CeO₂ and CeO₃ under oxidizing and reducing environments, respectively, and (b) the ease of formation of labile oxygen vacancies[18]. The selectivity of pure CeO₂ in oxidation reactions is poor, but the redox chemistry of doped cerium oxides is sensitive to crystal structure defects [19]. CeO₂ crystallizes in the fluorite structure in which each cerium ion is coordinated to eight oxygen neighbours, making CeO₂ more stable and the reduction of Ce (IV) to Ce (III) is unfavourable. To tackle this problem, one of the best approaches is replacing of another metal/metal oxide into the ceria lattice, thus facilitating the formation of composite oxides. The replacement of cerium ions by cations of different size and/or charge modifies ionic mobility within the lattice resulting in the formation of a defective fluorite structured solid solution. Such modifications in the structure of ceria impart new properties to the catalysts, such as density, ionic conductivity and lattice parameters, improved thermal stability, high catalytic activity [20,21]. Inspired by the unique and promising characteristics of ceria-based mixed oxides and solid solutions for various applications, a systematic study was undertaken to understand the of CeO₂–SiO₂, CeO₂–TiO₂, CeO₂–ZrO₂ and CeO₂–Al₂O₃ mixed oxides as catalyst support. Since, SiO₂, TiO₂, Al₂O₃ and ZrO₂ are extensively employed as supports as well as catalysts for several applications [22, 23]. We undertook the present investigation to stabilize CeO₂ by these oxides. The selected Support materials not only provide high reactive surface area, thermal stability, higher dispersion of nano-metallic particles and longer catalyst life but also act as oxygen exchangers and act as a sponge to capture and release oxygen molecules as per need during the reaction. Herein we report the aerobic oxidative dehydrogenation of amines to imines catalysed by gold supported on ceria based mixed oxides.

2. Materials and Methods

2.1. Catalyst preparation:

The first step is co-precipitation of metal oxide from metal nitrate solutions. A solution containing Ce(NO₃)₃.6H₂O (9.314 g) and SiO(NO₃)₄ (3.6052 g) is treated with a solution of NH₄OH at constant pH 9.0 and room temperature with constant stirring using mechanical stirrer. The resulting precipitates were aged at the same temperature for 12 h, then filtered and washed until the removal of nitrate ions. The washed precipitate is dried at 100 °C and calcined under air at 500 °C for 5 h. The support prepared in this way is Ceria-Silica and is denoted as CS. Similarly we prepared Ceria-titania (CT) using solution containing 8.6112 g Ce(NO₃)₃.6H₂O and 3.7276 g TiO(NO₃)₄. Ceria-Zirconia (CZ) by using 7.3014 g Ce(NO₃)₃.6H₂O and 3.915 g ZrO(NO₃)₂ and Ceria-Alumina (CA) by using 7.9169 g Ce(NO₃)₃.6H₂O and 6.8432 g Al(NO₃)₃.9H₂O following the procedure similar to that for the preparation of Ceria-Silica. All solid supports are in the 1:1 molar ratio for CeO₂:M₂O₃.

To prepare the catalysts Au/CS, Au/CZ, Au/CT and Au/CA, aqueous solution of HAuCl₄.3H₂O was precipitated by adding 1N aqueous NaOH at constant pH and temperature upon mixed metal oxide preliminary suspended in water by mechanical stirring. The resulting precipitate was aged at room temperature for 12 h, then filtered and washed carefully until complete elimination of
Cl− ions as detected by using silver nitrate as precipitating agent. The sample was dried at 80 °C and then calcined at 200 °C. The samples contain Au/CS=1.01, Au/CZ=1.15, Au/CT=1.79 and Au/CA=1.34 Wt% gold as determined by SEM EDEX method using instrument SEM Hitachi-S520, Japan; Oxford Link ISIS-300 UK with instrument operated at 98 eV resolution. The results were further confirmed by fluorescence X-ray crystallography using D8 – Advance, Bruker, Germany Multi Res-Vac34 method. Same method was used to determine Ce and the other metals which were found to be Ce=41.43 %and Si=20.32 % in Au/CS, Ce=42.27 % and Zr=18.87 % in Au/CZ, Ce=29.56 %and Ti=27.31% in Au/CT, Ce=37.48 %and Al=24.20% in Au/CA. The other constituents found were oxygen and minor impurities of Na, Ca, V, Rb and Th etc. The N2 BET surface areas of the powders were determined by N2 adsorption at liquid nitrogen temperature using a Micromeritics Gemini 2360 instrument. Prior to analysis, samples were oven dried at 120 °C for 12 h and flushed with Argon gas for 2 h. The surface area thus found was 107 m²/g for Au/CS, 101 m²/g for Au/CA, 98 m²/g for Au/CT and 87 m²/g for Au/CZ.

2.2. Catalytic Tests

After establishing the indispensable nature of the catalyst the conversion of pyrrolidine was chosen as a model reaction to explore the practicability of the proposed methodology (Scheme 1). The reactants used for the present work were of Analytical grade obtained from Fluka chemicals and were used as received. The reactions were carried out in two neck glass reactor fitted with a condenser in one neck and other neck was closed by a rubber septum through which a syringe needle was passed to supply oxygen (O2) from oxygen balloon. Reactant (0.1 mmol) was dissolved in toluene (2 mL) in the reactor to which 0.05 g of catalyst was added. Molecular oxygen was passed through syringe needle and the mixture was kept stirring using magnetic stirrer for 35 h at 100 °C.

The reaction mass was filtered to remove the catalyst powder. Products of these reactions were identified by comparing their mass spectra and GC retention times with those of authentic samples. Yields were determined by GC using authentic samples of the products as calibrants.

3. Results and Discussion

In this work, we selected conversion of secondary amines to imines as our model reaction to study the activity of ceria based mixed oxide supported nano-gold catalyst for selective oxidative dehydrogenation using molecular oxygen. After catalyst optimization it was found that Au/CT is most active catalyst among the asprepared catalyst so this catalyst was tried for different substrates (Table 1.). The catalyst recyclability was tested up to five reaction cycles and it was found that there is negligible loss in catalytic activity as well as selectivity even up to 5 cycle (Fig. 1). It was also found that loss of catalytic activity can be regained by washing the catalyst with piranha [24] (3:1 concentrated sulfuric acid: 30% hydrogen peroxide solution). On correlating the catalytic activity with the characterization of the catalysts (table 2.), it is observed that the surface area of the catalysts is not in agreement with catalytic activity but the gold content of the catalysts shows a close concurrence. The increase in catalytic activity with increase in gold content may be due to the increase in the number of active sites. The effect of gold

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Catalyst</th>
<th>Gold content (Wt %)</th>
<th>Ce Content (Wt %)</th>
<th>M Content (Wt %)</th>
<th>catalyst surface area (m²/g)</th>
<th>% Product Yield</th>
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<tbody>
<tr>
<td>1</td>
<td>Au/CS</td>
<td>1.01</td>
<td>41.43</td>
<td>20.32</td>
<td>107</td>
<td>86</td>
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<tr>
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<td>Au/CA</td>
<td>1.34</td>
<td>37.48</td>
<td>24.20</td>
<td>101</td>
<td>88</td>
</tr>
<tr>
<td>3</td>
<td>Au/CT</td>
<td>1.79</td>
<td>29.56</td>
<td>27.31</td>
<td>98</td>
<td>89</td>
</tr>
<tr>
<td>4</td>
<td>Au/CZ</td>
<td>1.15</td>
<td>42.27</td>
<td>18.87</td>
<td>87</td>
<td>80</td>
</tr>
</tbody>
</table>

Note: Reaction conditions: amine, 2 mL of solvent, catalyst powder (50 mg), O2 (1000 mL at 1 atm.) at 100 °C for 35 h in toluene
particle size is considered to be an important parameter controlling catalyst activity and it is well known that small gold particle size is highly active for selective oxidation because of their much larger surface area to mass ratio. But particle size increases with increasing Au content from 3% wt. onwards, because below 3% wt gold particles are highly dispersed and above this loading cluster size increase and makes only fewer sites available for catalytic activity [25]. In our case the most active catalyst Au/CT has highest gold content containing only 1.79 wt. % which is much lesser than limiting content (3 wt. %) among all the as-prepared catalysts, so there is no aggregation of gold nanoparticles.

On EDEX and BET surface analysis it is observed Au/CS has highest and Au/CZ has the lowest surface area among the as-prepared catalysts, the metal deposition is maximum in Au/CT and minimum in Au/CS. Au/CZ although has higher gold content than Au/CS but has lesser activity than latter. Making a look at CeO$_2$:MO$_2$ ratio it is clear that among all the four supports (or catalysts as a whole) the ratio is lowest for CT (Au/CT) and highest for CZ (Au/CZ), thus it can be said that the catalytic activity is inversely proportional to the CeO$_2$:MO$_2$ ratio. Ceria-titania combination shows higher catalytic activity among all the four catalysts, which may be due to the lower CeO$_2$:MO$_2$ ratio, the best fit of Ti$^{4+}$ into the CeO$_2$ lattice (Ti=1.3 Å and Ce=1.1 Å) and high dispersion capacity of titania [15]. From the above discussion it can be proposed that the catalytic activity of supported gold is affected by CeO$_2$:MO$_2$ ratio of the catalyst support and its composition also. This may be due to growth of crystalline size of mixed oxide with the increase in ceria content with the result a decrease in metal dispersion on the support, which leads to increase in Au cluster size, hence decrease in surface area of the active component and with the result decrease in catalytic activity [26]. It is reported that Au foil weakly chemisorbs CO but is unable to dissociate or molecularly adsorb H$_2$ and O$_2$, however, supported gold nanoparticles adsorb or react with O$_2$ [27] and it is also believed that there are two important factors for the activity of gold nanocluster catalysts, cluster size and a support [28]. Cluster size affects the activity of the catalysts by both surface area and quantum size effect [29]. Nano-gold 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 (Au$^0$, Au$^{1+}$ and Au$^{3+}$) and electronic properties of gold particles and, consequently, control the catalytic performance (both the activity and selectivity) of the supported nano-gold catalysts in the reactions [30].

<table>
<thead>
<tr>
<th>s. no</th>
<th>substrate</th>
<th>product</th>
<th>% yield</th>
<th>% selectivity</th>
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<td>1</td>
<td>(\text{NH}_2\text{NH})</td>
<td>(\text{N} - \text{N})</td>
<td>89</td>
<td>&gt;99</td>
</tr>
<tr>
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<td>(\text{NH}_2\text{NH})</td>
<td>(\text{N} - \text{N})</td>
<td>85</td>
<td>&gt;99</td>
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<tr>
<td>3</td>
<td>(\text{N} - \text{H})</td>
<td>(\text{N} - \text{H})</td>
<td>78</td>
<td>&gt;99</td>
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<td>(\text{N} - \text{H})</td>
<td>(\text{N} - \text{H})</td>
<td>71</td>
<td>&gt;99</td>
</tr>
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</table>

Note: Reaction conditions: amine, 2 mL of solvent, catalyst powder (50 mg), O$_2$ (1000 mL at 1 atm.) at 100 °C for 35 h in toluene

**Figure 1:** Reaction cycles of Au/CT catalyst for the conversion of pyrrolidine to 3,4-dihydro-5-(pyrrolidin-1-yl)-2H-pyrrole
4. Conclusion

It can be concluded that nano-gold supported on CeO$_2$-M$_n$O$_a$ is a suitable, efficient, and selective and robust catalyst for the oxidative dehydrogenation of secondary amines to imines in aerobic conditions. It is also observed that support effects the deposition of metal and also the activity of the catalyst. This process is very selective, productive and the catalyst though is costly but it is recyclable which compensates the cost so we can propose that the process is economically beneficial.

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References


