STRUCTURE, CHARACTERIZATION AND APPLICATION OF NI HYDROTALCITE AS SOLID BASE CATALYSTS FOR ORGANIC TRANSFORMATIONS

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(Received: December 29, 2009 - Accepted: January 26, 2011)

ABSTRACT

Ni hydrotalcites with different kinds of metals, ions exhibit considerable activity for organic transformations. Various ratios of Ni-Al hydrotalcites are prepared by using different compositions which are characterized by XRD, IR and TPR analysis. Carbonate in the brucite sheet is active for selective oxidation reactions. Upon calcination the dehydroxylation of the hydrotalcite and concurrent decomposition of the carbonate ions produce mixed oxides which exhibit activity for variety of organic transformations.

Keywords: Hydrotalcite, LDHs, mixed oxides, structure.

1. INTRODUCTION

Heterogeneous catalysts have many advantages such as easy separation and reusability. The efficient catalyst have many economic benefits such as ease of catalyst recovery, low cost, recycling, zero emission of pollutants, longer life time. Till 1970 the research into the catalysis, and hydrotalcites were followed in parallel paths then the first patent which was referred to as hydrotalcite was prepared for the use in hydrogenation reactions [1].

Hydrotalcite, which is a mineral white crushed into fine powder and first discovered in Sweden around 1842 as a hydroxy carbonate of magnesium and aluminium which occurs in nature in foliated and contorted plates. The first formula of hydrotalcite was presented by Professor Mannase, at the University of Florence [2].

The hydrotalcites are classified as Cationic or Smectite type in which the layered structures are arranged in sheets as brucite type Mg(OH)\(_2\) and Al(OH)\(_3\), octahedral separated by charge balancing anions, water and these occupy the interlayer spaces. The sheets then stack (in the crystallographic c direction) to give a characteristic layered material. In many samples of hydrotalcite, few of the M\(^{2+}\) species are substituted by M\(^{3+}\) species, which results into the layers carrying the residual positive charge. Crystals of hydrotalcite which mostly consists of micron sized stacks of hexagonal platelets morphologically. However, fibrous and extended sheet-like morphologies is reported depending upon the conditions of synthesis [4-5]. Thermal calculation of these materials results in the formation of non-stoichiometric mixed metal-oxides with characteristic properties, which are extremely active catalysts for many important transformations. Various catalytic applications of hydrotalcite are used for steam reforming, methanol and higher alcohol synthesis [6], selective oxidation of alcohols to aldehydes [7], and reduction of aldehydes to alcohols [8], epoxide ring opening [9], epoxidation [10], Aldol condensation [11], cyanoethylation [12], Michael reaction [13], and transterification [14].

Pinnavaia [15] and co-workers reported excellent work on intercalation of polynuclearmetallates as anion pillars and exploited them in selective oxidation reactions [16]. During the past decade strong impetus was given to modify these materials for the synthesis of fine chemicals, exploring both basic and redox properties. Recently Choudary et al [17] reported oxidative bromination of bisphenol-A excellent activity for some intercalated hydrotalcite for asymmetric dihydroxylation of various aliphatic and aromatic olefins [18] using NMO (N-methylmorpholine oxo) as co-oxidant. The maximum amount of water present in the interlayer can be calculated on the basis of sites present in the close packed configuration of oxygen atoms, as discussed here Kannan and co-workers reported intercalation of metallophthallocyanine tetrasulfonate in the interlayer of Mg-Al hydrotalcite which is synthesized by different routes. Many variations in compositions are reported for hydrotalcites. Hydrotalcite catalysts are prepared by different co-precipitation methods depending upon the composition of the M\(^{2+}\) and M\(^{3+}\) [20]. Choudary et al [7] reported Ni-Al hydrotalcite is prepared by the above said methods. Ni-Al hydrotalcite with varied composition of Ni-Al hydrotalcites (Ni:Al=2:1 (cat. A), 2.5:1, 3:1) were prepared with co precipitation technique by employing NaOH/Na\(_2\)CO\(_3\) as described in the literature [20-22].

3. RESULTS AND DISCUSSION

3.1. Catalysts Characterization:

XRD of Hydrotalcites:

The XRD for LDHs samples such as carbonates which have one unit hexagonal cell as shown in Fig. 1. The peak at 11.5° 2\(\theta\) (d spacing = 7.7 Å) of hydrotalcite carbonate was attributed to the reflections from the (003) family of crystallographic planes. These planes are spaced one-third per unit cell distance apart and these corresponds to the interlayer distance [23] studied by Carlino. The d spacing is known as the interlayer spacing which is formed from the metal hydroxide sheet (approximately 4.8 Å) and the gallery region is around 2.8 Å contains the intercalated anions. This gallery between the metal hydroxide layers is known as the interlayer distance or gallery height. The XRD samples
of calcined at 723 K mainly shows the presence of mixed cubic oxide phases, which is obtained by the loss of water and carbon dioxide from the anionic interlayer of hydrotalcite precursors. Ni-Al 2:1, 2.5:1, 3:1 hydrotalcite shows the characteristic crystalline pattern explained above which shows excellent activity for oxidation of alcohols with Ni-Al HT with molecular oxygen [7]. The used catalysts retained similar hydrotalcite XRD pattern. The XRD of the Ni-Al hydrotalcite calcined (Cat A) shows formation of NiO phase only with Al(III) inside. When the catalysts were rehydrated for 2 days at room temperature, the calcined sample could not be restructured into hydrotalcite [7-8]. These results are in good agreement with the reported data that restructured calcined Ni-Al hydrotalcite requires stringent conditions [16-17,24].

Figure 1. XRD powder patterns of samples. (a) Ni-Al-CO$_3$ as-synthesised, (b) calcined Ni-Al hydrotalcite catalyst (cat A) and (c) used Cat A [8].

**Infrared Spectroscopy of Hydrotalcites:**

The infrared spectra for the hydrotalcites hydroxy stretching vibrations of Ni-Al hydrotalcite [Cat. A (Ni:Al 2:1)] and Ni-Al hydrotalcite (Ni:Al 3:1) are shown in Fig. 2. The IR spectra of MgAlCO$_3$ hydrotalcite, with different M(II) cations and Ni-Al hydrotalcites showed the absorption at 3500-3600 cm$^{-1}$, which is attributed to the H-bonding stretching vibrations of the OH group that is present in the brucite like sheets. Mostly the shifting of this bond depends upon the $x$ for Mg(OH)$_2$. When $X=0$ the absorption of this band shifts to higher frequency of 3700 cm$^{-1}$. Cavani et al [25] reported that as the (M$^{2+}$/M$^{3+}$) ratio increases the hydrogen stretching & bending corroborated with the changes in the layer spacing. A shoulder is present around 3000cm$^{-1}$, is mainly attributed to hydrogen bonding between H$_2$O and the anion in the interlayer [16,25] and H$_2$O bending vibration which occurs at 1600 cm$^{-1}$. In most hydrotalcites [25] the three bands appears at 1350-1380cm$^{-1}$ (v$_s$), 850-880 cm$^{-1}$ (v$_s$), and 670-690cm$^{-1}$. The IR spectra of various compositions of Ni-Al hydrotalcite shows that the vibrational stretching frequency of the hydrogen atom in hydroxide group of Cat A with 2:1 Ni-Al ratio appears at lower wavelength 3420 cm$^{-1}$ and has a smaller half width than the other compositions which indicates there is a more ordered cation distribution in the former [7-8].

**Figure 2.** Hydroxy stretching vibrations of a) Ni-Al hydrotalcite, Cat. A (Ni:Al 2:1) and b) Ni-Al hydrotalcite (Ni:Al 3:1) [7].

**TPR Characterization:**

TPR studies are used for the characterization of the heterogeneous catalytic systems and are shown in Fig. 3. The TPR profile of hydrotalcite indicates that free NiO is reduced at lower temperature and the NiO in association with oxide of aluminium is reduced at higher temperatures and ammonia display both free NiO and NiO in association with oxide of aluminium nickel aluminium hydrotalcite displays a shoulder at lower temperature that grows on free nickel oxide content. These TPR results are in good agreement with the results reported by Choudary et al [7]. The TPR samples of Ni/γ-Al$_2$O$_3$ indicate the presence of free NiO only. The catalysts are prepared with ammonia method, displays both free NiO and NiO in association with aluminium oxide. The TPR for nickel containing hydrotalcite was studied for Mg-Ni of varying ratios, showed two peaks of H$_2$ consumption. The first peak around 570 K corresponds to the release of NO$_3$ anions as NO$_2$ and the subsequent reduction of NO and N$_2$O. The second peak with maxima at 705, 920 and 1000 K for respective samples to the reduction of NiO particles. When Mg content is increased the reduction of the nickel oxide decreases it can be compared with the decrease of the NiO crystal size. This behaviour is attributed to the formation of Ni aluminate band of nickel spinel type and decreasing the size the crystallite and hence it hinders their reducibility [26-31]. The excellent results of these catalysts from characterization studies guide us to investigate for high activity of these catalysts Ni-Al 2:1 for the bromination of β-ketoester at α-positon with N-bromosuccinimide as brominating agent. The conversions were 65% with high selectivity. Whereas, with the Ni-Al 2.5:1 the conv was 60% and with 3:1 the conv was 57%. The less conv activity of Ni-Al 2.5:1 and 3:1 were due to high nickel content [32] present, but with high selectivity retained.
Figure 3 TPR profiles plotted as amount of hydrogen consumed (x is arbitrary units) versus temperatures [°C] of a) Ni-Al hydrotalcite (cat A) b) Ni-Al hydrotalcite (Ni: Al 2.5:1), c) Ni-Al hydrotalcite (Ni: Al 3:1) d) Ni-γ-Al$_2$O$_3$ 2% e) Ni-γ-Al$_2$O$_3$ 5% f) Ni-γ-Al$_2$O$_3$ 10%, g) Ni-Al Cat B [7].

4. CONCLUSIONS

In conclusion Ni hydrotalcites have been characterized by XRD, IR and TPR results which confirms that nickel hydrotalcites are active catalysts for oxidation of alcohols, alkyl aromatics, epoxidation of olefins, sulphur oxidation and reduction, reactions, our group is involved in studying organic transformations such as Aldol, Micheal, Knoevenegal, oxidation of alcohols, dihydroxylation of olefins, bromination of bis-phenol-A with Ni containing hydrotalcites. The excellent results of these catalysts from characterization studies guide us to investigate the high activity in bromination of β-ketoester at α-position for organic transformations.

ACKNOWLEDGEMENTS

The authors extend their appreciation to the deanship of scientific research at king saud university for funding the work through the research group project No RGP-VPP-089.

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