Recent Progress of Hydrogenation and Hydrogenolysis Catalysts Derived from Layered Double Hydroxides

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Abstract: Layered double hydroxides (LDHs), also known as hydrotalcite-like compounds, are widely used in many fields due to their unique structural advantages. Based on LDHs, a wide range of metal catalysts could be synthesized with high metal dispersion, tunable acid-base properties, facile but flexible preparation methods, strong metal-support interaction, and thermal stability. Owing to these outstanding advantages, LDH-derived materials manifest great potential as catalysts, particularly in hydrogenation and hydrogenolysis reactions. More than 200 papers published in the past five years in this field clearly indicated the rapid development of these materials. In this respect, it is imperative and essential to provide a timely review to summarize the current progress and motivate greater research effort on hydrogenation and hydrogenolysis catalysts derived from LDHs. In this review, the applications of LDH-derived materials as heterogeneous catalysts in various hydrogenation and hydrogenolysis reactions were comprehensively discussed. Hydrogenation of unsaturated chemical bonds, hydrodeoxygenation of oxygenated compounds, hydrogenolysis of carbon–carbon bonds and hydrogenation of nitrites and nitriles were described. This review demonstrates the extraordinary potentials of LDH-derived catalysts in hydrogenation and hydrogenolysis reactions, and it is undoubted that LDH-derived catalysts will play an even more significant role in the foreseeable future.

Keywords: layered double hydroxides; hydrotalcite; hydrogenation; hydrogenolysis; catalysis

1. Introduction

Hydrogenation is one the most fundamental processes in the chemical industry with diverse applications. In hydrogenation, atomic hydrogen is added to unsaturated chemical bonds in the presence of homogeneous or heterogeneous catalysts. Hydrogenolysis is another important type of reaction where a carbon–carbon bond or carbon–heteroatom bond is cleaved by hydrogen. In the past century, hydrogenation and hydrogenolysis are particularly important in the petrochemical industry to saturate alkenes or aromatics, remove sulfur or nitrogen atoms, increase fuel stability and decrease toxicity [1]. Recently, hydrogenation and hydrogenolysis attracted extensive attention for new objectives. Biomass utilization [2,3] and CO₂ hydrogenation [4,5], which are crucial processes for a sustainable human society, also heavily rely on the technological development of hydrogenation and hydrogenolysis. Using renewable H₂ produced from eco-friendly processes [6,7], hydrogenation and hydrogenolysis in biomass upgrade and CO₂ hydrogenation could greatly reduce the dependency on fossil fuels. In most industrial cases, hydrogenation or hydrogenolysis are catalyzed by supported metal catalysts based on transition metals such as Ni, Cu, Pd, or Pt [2,8,9].

Layered double hydroxides (LDHs) are layered inorganic materials consisting of positively charged brucite Mg(OH)₂-like layers and interlayer charge-compensating anions, as
shown in Figure 1 [10]. A typical type of LDHs is hydrotalcite (HT), a natural mineral with chemical composition of Mg$_6$Al$_2$(OH)$_{16}$CO$_3$·4H$_2$O [11]. LDHs and natural hydrotalcite share similar structure, in which metal cations can be replaced by divalent and trivalent metal ions with similar atomic radii in the same crystal, and interlayer anions can be replaced by other intercalating anions. LDHs, also called hydrotalcite-like compounds, could be expressed with the general formula $[M^{2+}_{(1−x)}M^{3+}_x(OH)_{2}]^{x+}·[(A^{n−})_{x/y−2}H_2O]^{y−}$ [12]. Common $M^{2+}$ ions include Mg$^{2+}$, Cu$^{2+}$, Zn$^{2+}$ and Ce$^{2+}$, while common $M^{3+}$ include Al$^{3+}$, Fe$^{3+}$, and Cr$^{3+}$. These cations can be atomically and uniformly dispersed in the hydrotalcite layer. The formed cation layers lack interlayer interaction and have a negative net charge, which must be balanced by the inserted anions, and water molecules might also enter the interlayer region [13]. Interlayer anions are exchangeable, such as CO$_3^{2−}$, NO$_3^{−}$, SO$_4^{2−}$, Cl$^{−}$, F$^{−}$, acetate and salicylate. The $x$ value in the general formula represents the molar ratio of $M^{2+}/(M^{2+} + M^{3+})$, and the range of its value has great influence on the composition and structure of hydrotalcite materials [10]. Both the magnitude of the charge density on the laminate and the extent to which the homocrystalline substitution of metal cations occurs depend on the $x$ value, and a genuine LDH phase is usually found only at $0.2 < x < 0.4$ [14,15].

![Figure 1. A Schematic illustration of LDH structure.](image)

The unique structure, tunability of the layer and interlayer element composition, delamination property, structural topological transformations, and the confinement effect of LDHs provide LDH-derived materials great application potentials in many fields [14,16]. In heterogeneous catalysis, LDH can be applied as catalyst itself, catalyst support or catalyst precursor in many reactions such as C-C coupling, N-arylation, oxidation, hydrogenation, hydrogenolysis, etc. Although several review articles on catalytic applications of LDHs have been published [12,17–21], reviews focusing on hydrogenation and hydrogenolysis are still scarce to the best of the authors’ knowledge. Furthermore, in the past few years, a large number of research articles on hydrogenation and hydrogenolysis over LDH-derived catalysts were published, demonstrating the significant development in the field. Therefore, we believe it is timely to provide a comprehensive and in-depth summary on the latest progress of LDH-derived catalysts for hydrogenation and hydrogenolysis. This review will focus on the applications of LDH-derived materials as hydrogenation and hydrogenolysis catalysts, highlighting physiochemical properties, structural properties and reactivity advantages of LDH-derived catalysts. Thermocatalytic processes catalyzed by heterogeneous catalysts will be the emphasis of this review, while electrocatalytic or photocatalytic research will not be discussed due to their distinct nature in mechanisms.
2. Type of LDH-Derived Catalysts for Hydrogenation and Hydrogenolysis

Generally, three types of LDH-derived catalysts are prepared and investigated: LDH-supported catalysts, LDH-derived mixed metal oxides (MMOs), and LDH-derived intermetallic compounds (IMCs). Table 1 shows several examples of different types of LDH-derived catalysts and their reported physiochemical properties. It could be noticed in the table that high specific surface area and small particle size could be readily achieved with catalysts from LDHs. This section will describe in detail the general advantages and synthesis protocols of these three types of LDH-derived materials.

| Catalyst          | Preparation Method                  | Surface Area (m²/g) | Particle Size (nm) | Other Properties                                                                 | Ref.  |
|-------------------|-------------------------------------|---------------------|--------------------|----------------------------------------------------------------------------------|-------|
| PdAg/ZnTi-LDH     | Coprecipitation and photochemical reduction | 131                 | 5                  | Pore volume 2.63 cm³·g⁻¹; mean pore size 80.3 nm                                  | [22]  |
| Pt/CoAl-MMO       | Coprecipitation and reduction–deposition | 87                  | 4–5                | Pore volume 0.29 cm³·g⁻¹; mean pore diameter 3.83 nm                             | [23]  |
| CuMgAl-MMO        | Urea hydrolysis                      | 213                 | –                  | Pore volume 0.95 cm³·g⁻¹                                                          | [24]  |
| CuCoAl-MMO        | Coprecipitation                      | 92.9                | 3.1                | Pore volume 0.69 cm³·g⁻¹                                                          | [25]  |
| NiMoAl-MMO        | Coprecipitation and ion exchange     | 93                  | 4.6                | Pore volume 0.13 cm³·g⁻¹; total acid sites 2.077 mmol·g⁻¹                         | [26]  |
| NiIn-IMC          | Co-precipitation                     | 126.4               | 5.8                | –                                                                                | [27]  |
| NiMo-IMC          | In situ co-reduction                 | 189.2               | 18.6               | –                                                                                | [28]  |

2.1. LDH-Supported Catalysts

Because heterogeneous catalysts for hydrogenation and hydrogenolysis usually require the participation of metallic sites, pristine LDHs could not be used directly as hydrogenation and hydrogenolysis catalysts. Nevertheless, LDHs act as perfect support materials due to intrinsic structural superiorities. High specific surface area, hierarchical pore structure, abundant surface defects, strong metal-support interaction, surface acid-base sites with tunable strength or concentration, or inherent confinement effects have been reported for LDHs [14,20].

Major synthesis methods of LDHs include coprecipitation [29], urea hydrolysis, hydrothermal synthesis [30] and the sol-gel method [15]. Different synthesis methods directly affect the physical and chemical properties of the obtained LDH materials [31–33]. The most facile and common method is coprecipitation. By coprecipitation, crystallinity, particle size distribution, and stability of LDHs can be precisely controlled by adjusting the synthesis parameters such as pH, temperature, aging time, mixing rate, cation ratio and solution concentration [31,32]. Coprecipitation can be combined with the anion exchange method to modulate the structure and function of LDH-based materials by inserting ions or molecules between layers while keeping the basic structure unchanged, modifying the physiochemical properties or the surface properties of the material.

2.2. LDH-Derived Mixed Metal Oxide Catalysts

Mixed metal oxides (MMOs), which could be synthesized by the heat treatment of LDHs, showed high catalytic performance and received broad attention in hydrogenation and hydrogenolysis. LDH-derived MMOs inherit LDH characteristics such as controlled composition, high specific surface area, and uniform morphology. In addition, by converting metal hydroxides into metal oxides, highly dispersed metal oxides particles were generated, which could act as acid or base sites. The nature, intensity and concentration of these acid/base sites on MMOs could be tuned by controlling the type and molar ratio of metal cations in the precursors, preparation method, calcination temperature and
type of interlayer anions [20,34,35]. The high surface area and tunable acid/base property, as well as other advantages, make LDH-derived MMOs an ideal catalyst support material compared to the conventional metal oxide support. MMOs could also be used independently as hydrogenation/hydrogenolysis catalysts. For MMOs with hydrogenation-active metals such as Ni or Cu, the reduction of MMOs could generate highly dispersed metallic sites in situ which are catalytically active for hydrogenation/hydrogenolysis. Compared with supported metal catalysts prepared by impregnation, reduced MMOs as hydrogenation/hydrogenolysis catalysts are often more reactive, due to the high metal dispersion, abundant acid/base sites and strong metal–support interaction, as shown later in this review.

The calcination of LDHs to generate MMOs is a topological transformation process. During thermal transformation three temperature domains of LDH structure changes can be observed. The first structural change at around 190 °C corresponds to the dehydration of loosely bonded interlayer water, accompanied by the decrease in the layer spacing, while the layered structure is retained [36]. The second temperature domain in the range of 200–400 °C indicated dehydroxylation and the collapse of the layered structure of LDHs, in situ forming highly dispersed metal oxides [37]. The third domain in the range of 500–1000 °C is ascribed to the formation of sintered metal particles and spinel phases [36,38]. The structure and function of MMOs can be controlled by changing the topological transformation parameters such as heating rate, calcination temperature and calcination atmosphere of LDH precursors during the thermal treatment, leading to the modification of the catalytic performance of MMOs [10,17,34,39,40].

2.3. Intermetallic Compound Catalysts

Intermetallic compounds (IMCs), also called ordered alloy, is a special type of alloy composed of two (or more) metal elements with specific crystal structure and atomic composition. IMCs can be expressed as $A_xB_y$, where $x$ and $y$ are small integers, usually 1, 2 and 3 [41,42]. Compared with conventional alloys, IMCs exhibit completely or partially ordered surface structure. Figure 2 showed the structure of NiSb and NiBi IMCs catalysts prepared by Wei’s group [43]. It could be noticed that high metal dispersion and uniform alloy composition were obtained with IMCs, which were important for catalytic reactivity. Due to the unique electronic and geometric properties, IMCs promote the activated adsorption of substrate molecules with specific patterns or configurations, thus achieving improved catalyst activity, selectivity and stability [41,44,45].

![Figure 2](image-url)
The preparation of IMCs can be categorized into three methods, controlled colloid synthesis, inorganic capsule synthesis, and layered double hydroxide synthesis [42]. The synthetic method based on LDHs has particular advantages. The composition flexibility of LDH precursors enable the possibility of synthesizing a variety of IMC catalysts [45,46]. Metal elements of LDH layers have a high degree of dispersion at the atomic level, which is conducive to the formation of metal catalysts with high metal dispersion after the calcination-reduction process. The preparation of IMC nanocrystals using LDH precursors can be achieved by two methods, endogenous method or exogenous method. For the endogenous method, two metal components of IMCs (e.g., Ni, Co, Cu, Ga) are introduced simultaneously into the host layer of LDHs, followed by the calcination and reduction of LDH precursors, affording IMC catalysts. For noble metal elements (Au, Rh, Pd, Ir, Pt, etc.) or heavy metal elements (Sb, Sn, Bi, Pb, etc.) which are difficult or impossible to introduce into the LDH layer, the exogenous method can be adopted, in which noble metal or heavy metal salts are mixed with LDH precursors by impregnation, followed by calcination and reduction to obtain IMC catalysts.

3. Hydrogenation of Carbon-Oxygen Unsaturated Bonds

3.1. Hydrogenation of Ketones and Aldehydes

The hydrogenation of carbonyl groups in ketones and aldehydes to form hydroxy groups is an important process particularly in synthetic chemistry. This section will review research work on ketone or aldehyde hydrogenation catalyzed by LDH-derived catalysts. The selective hydrogenation of \( \alpha - \beta \) unsaturated aldehydes including crotonaldehyde, citral and cinnamaldehyde will also be discussed. The hydrogenation of furfural, a highly important furan aldehyde derived from hemicellulose, will be reviewed in the next section separately.

For hydrogenation over heterogeneous catalysts, metal dispersion is crucial because high dispersion would naturally mean more active sites for reactions to take place. Non-precious metal catalysts such as Ni, Cu and Co could be synthesized by the calcination of corresponding LDH precursors prepared by coprecipitation. Because \( M^{2+} \) and \( M^{3+} \) ions are uniformly distributed in the brucite layers, the calcination of LDHs could in situ generate highly dispersed metal particles and more catalytic active sites for hydrogenation. For instance, Dragoi’s group have shown that a reduced CuMgAl catalyst synthesized from LDH precursors exhibited particle sizes of 2.6 to 6.5 nm and high activity in cinnamaldehyde hydrogenation to cinnamyl alcohol [40]. Another study using CuMgAl-MMO of 1.4 to 2.4 nm metal particle size for benzyl aldehyde hydrogenation, obtained 85–93% selectivity of benzyl alcohol [47].

As shown in Table 2, precious metals are particularly effective in selective hydrogenation of unsaturated aldehydes. For precious metal catalysts such as Pt, Pd or Ru, using LDHs or LDH-derived MMOs as support could also generate catalysts with high precious metal dispersion. Many examples were found in the literature reporting metal particle sizes less than 5 nm using the conventional impregnation method on LDHs/MMOs owing to the abundant surface basicity to anchor metal species [48], such as Ru/MgAl-HT [49], Pd/MgAl-MMO [50] and Au/MgAl-MMO [51] for aldehyde/ketone hydrogenation. Notably, due to the special interlayer galleries of LDHs, an alternative way to load metals to the support other than impregnation is by intercalating metal-containing anions in LDHs by the ion exchange method. With this method, researchers were able to obtain a Pt/MgAl-HT catalyst with an average diameter of Pt nanoparticle being only 1.9 nm using PtCl\(_4^{2-}\) [52]. Generally, as the catalyst particle size decreases, the number of low coordination sites such as the edges and corners of the metal increases, providing more active sites for the reaction and thus enhancing the catalytic activity of the catalyst [23,27,53].
Table 2. Catalytic performances of LDH-derived catalysts for hydrogenation of ketones and aldehydes.

| Catalyst | Substrate | Reaction Conditions | Conversion (%) | Product Selectivity (%) | Ref. |
|----------|-----------|--------------------|----------------|------------------------|------|
| Pt/ZnSnAl/C | 2-Pentenal | 80 °C; 3.0 MPa H₂ | 28.5 | 2–Pentenol, 92.0 | [54] |
| Cu/MgAl-HT | Benzaldehyde | 250 °C | 68 | | [47] |
| Ru/MgAl-HT | Benzaldehyde | 100 °C; 3.5 MPa H₂ | >63 | Cyclohexanemethanol, 91.9 | [49] |
| Au/Mg₂AlO | Crotonaldehyde | 120 °C; 0.93 MPa H₂ | 23.6 | Crotyl alcohol, 62 | [51] |
| Pt/MgAl-LDH | Cinnamaldehyde | 80 °C; 2 MPa H₂ | 92.6 | Cinnamyl alcohol, 75.5 | [52] |
| Pt/CoAl-MMO | Cinnamaldehyde | 70 °C; 2 MPa H₂ | 99.7 | Cinnamyl alcohol, 72.5 | [23] |
| Au/ZnAl | Cinnamaldehyde | 130 °C; 1.5 MPa H₂ | 100 | Cinnamyl alcohol, 95.7 | [53] |
| Pt/MgAl-LDH | Cinnamaldehyde | 60 °C; 1 MPa H₂ | 79.7 | Cinnamyl alcohol, 85.4 | [55] |
| PtGa/MgAlGa | Cinnamaldehyde | 70 °C; 3 MPa H₂ | 52.8 | Cinnamyl alcohol, 70.7 | [56] |
| Ir/MgAlFe | Cinnamaldehyde | 60 °C; 3 MPa H₂ | 94.4 | Cinnamyl alcohol, 79.1 | [57] |
| Pt/CoAl-LDH | Cinnamaldehyde | 70 °C; 3 MPa H₂ | 94.3 | Cinnamyl alcohol, 91.9 | [58] |
| CoGa-IMC | Cinnamaldehyde | 100 °C; 2 MPa H₂ | 100 | Cinnamyl alcohol, 96 | [59] |
| NiZnAl/C | Citral | 140 °C; 1 MPa H₂ | 100 | Citronellol, 92.3 | [60] |
| CoSn-IMC | Citral | 160 °C; 4.0 MPa H₂ | 100 | Citronellol, 67.6 | [61] |
| CuZnAl-MMO | Unsaturated aldehydes | 80 °C; 1.0 MPa H₂ | 99.8 | Allylic alcohol, 75.1 | [62] |
| NiBi-IMC | Unsaturated aldehydes | 100 °C; 2 MPa H₂ | >90 | Unsaturated alcohol, >93.2 | [43] |
| NiSn-IMC | Unsaturated aldehydes | 120–145 °C; 3 MPa H₂ | >56 | Unsaturated alcohol, >44 | [27] |
| Pt/MgCoAl | Unsaturated aldehydes | 80 °C; 2 MPa H₂ | >87 | Unsaturated alcohol, >80 | [63] |

Because metal particle size plays an important role in the reactivity of the catalyst, particularly for the selective hydrogenation of α–β unsaturated aldehydes, several methods were developed for tuning particle size in LDH-related catalysts. Xiang et al. prepared HT-supported Pt nanocrystal catalysts for the selective hydrogenation of cinnamaldehyde [55]. It is noticed in this research that the selectivity of cinnamyl alcohol was closely related to Pt particle size, while the size of the Pt nanocrystals could be finely tuned by controlling the amount of surfactant added. Another study provides an example of using different alcohols including ethylene glycol, methanol and ethanol for the reduction of Pt on MgAl-LDHs, generating different sizes of reduced Pt particles with distinct reactivity in cinnamaldehyde hydrogenation [64].

For LDH-derived catalysts, the fact that active metal particles were incorporated in the LDH or MMO matrix indicates strong metal-support interaction (SMSI) [65]. SMSI is a specific interaction effect between the support and the metal nanoparticles (NP). This interaction may lead to metal–metal bonding or the formation of intermetallic compounds in the catalyst [66], enhancing the stability of the metal particles in the catalyst. In addition, the topological transformation from LDHs to MMOs can be exploited to create SMSI by stabilizing the active metal particles in the oxide matrix, to prevent leaching and agglomeration [67–69]. Wang et al. reported a slight increase in particle size from 1.2 nm to 1.6 nm for the Pt catalyst supported on ZnSnAl-MMOs after 12 h of 2-pentenal hydrogenation reaction [54]. Metal–support interaction could also electronically and geometrically affect metal atoms, shifting the product distribution in selective hydrogenation [56,57,67]. For instance, the addition of ZnO into NiZnAl-MMOs could deactivate C=C adsorption on Ni sites, inducing increased citronellol selectivity in citral hydrogenation [60]. Oxygen vacancies, which play important roles in carbonyl hydrogenation, could also be readily generated by SMSI. Miao et al. reported the formation of oxygen vacancies at the metal–support interface of Pd/MgAl-LDHs [70] and Pt/CoAl-LDHs [58]. It is found that these oxygen vacancies are activation sites for C=O bonds, and significantly enhance the selectivity toward C=O hydrogenation over C=C bond hydrogenation. Although SMSI is important for catalysts, its actual mechanism on product selectivity is still complex and case-dependent. It is proposed in a study that SMSI could enhance the C=C bond adsorption on Pd sites on MgAl-LDHs, leading to an increased selectivity toward citronellal in citral hydrogenation [71], while another research on cinnamaldehyde hydrogenation over Pt/CoMgAl suggested that SMSI strengthened the carbonyl adsorption and inhibited C=C hydrogenation [63].
Wei’s groups synthesized a series of intermetallic compound (IMC) catalysts using LDH precursors, including CoGa, CoIn, NiBi, CoSn and NiIn, for selective hydrogenation of unsaturated aldehydes \[43,59,61\]. It could be concluded from their research that by forming IMC catalysts, the electron density of active metal (Ni or Co) was modified due to electron transfer, leading to a change in adsorption conformation or adsorption strength which eventually affected product selectivity. The promising results from IMCs, as well as the convenient preparation method using LDH precursors, suggest a promising prospect of IMCs in heterogeneous catalysis.

For $\alpha - \beta$ unsaturated aldehydes, total hydrogenation of both C=C bonds and C=O bonds will also decrease yields toward desired products, and in most cases selective hydrogenation is pursued rather than total hydrogenation. The tunable acid–base properties of LDHs or MMOs play an important role in product distribution of selective hydrogenation of $\alpha - \beta$ unsaturated aldehydes \[72,73\]. It is shown that CuZnAl-MMO catalysts with the most Lewis acid sites which adsorbed citral molecules via the carbonyl group showed the highest selectivity toward allylic alcohol products such as geraniol and nerol \[62\]. Another study on Pt/MgAl-MMO also suggested that catalysts with stronger acidity were more prone to unsaturated alcohol production \[74\]. It is deduced that Lewis acid sites, such as metal cations, are adsorption and activation sites for C=O bond, and thus the selectivity of C=O bond hydrogenation products is closely associated with Lewis acidity of catalysts. LDH-derived catalysts often exhibit a close vicinity to Lewis acid sites for carbonyl adsorption and metal sites for hydrogenation, inducing synergistic effects for efficient hydrogenation of carbonyl bond in aldehyde or ketone hydrogenation \[75\].

### 3.2. Hydrogenation of Furfural

Furfural (FAL) is an important biomass platform chemical bridging biomass feedstock and bio-derived chemicals with an annual production exceeding 652 kilotons \[76\]. FAL could be converted from lignocellulose and their chemical versatility implies plentiful potential applications in the production of renewable C$_5$ chemicals \[77\]. C=C double bond and C=O double bond in furfural molecules require selective hydrogenation capability from catalysts to yield desired products. The most prevalent product derived from furfural is furfuryl alcohol (FOL), as more than 65% of FAL produced worldwide was converted into FOL \[78\]. The hydrogenation of FAL to yield furfuryl alcohol (FOL) could be achieved by various metals \[77\]. Non-precious metal Cu- and Ni-based catalysts prepared from LDH precursors were studied for carbonyl group hydrogenation to produce furfuryl alcohol, such as CuAl \[79,80\], CuCr \[81,82\], NiAl \[83,84\], NiSn \[85\], and CuNiAl \[86,87\]. Advantages of preparing Cu or Ni catalysts from LDH precursors are the small metal particle size and high stability, due to the uniform atom distribution in LDH precursors. Yang et al. reported the NiSn intermetallic compound catalyst prepared from LDH precursors exhibited a monometallic Ni particle with a size of around 10 nm. The high dispersion and the electronic modification effect of Sn resulted in a 99% selectivity toward FOL from FAL at 100 $^\circ$C and 2 MPa, and a high stability after more than 30 h \[85\].

Considering that the C=O bond from the furfural molecule could interact with the basic site on the catalyst surface, metal catalysts with surface basicity were synthesized, with Mg being the most commonly used metal due to the facile preparation from MgAl-hydrotalcite. NiMgAl-MMOs synthesized by calcining hydrotalcite precursors displayed high reactivity for the hydrogenation of FAL to FOL, with the synergistic effect between metallic Ni species to dissociate H$_2$ and surface basic sites generated by Mg introduction to activate furfural via carbonyl groups \[88\]. Villaverde et al. prepared CuMgAl catalysts derived from HT-like phases possessed high copper dispersion as well as strong interactions between metallic copper and magnesium–aluminum support, resulting in better activity, selectivity and stability than impregnated Cu catalysts \[89,90\]. Since CuNiAl, NiMgAl, and CuMgAl all showed outstanding reactivity in furfural hydrogenation, it is reasonable that CuNiMgAl would also display desirable catalytic properties \[91,92\], and it is demonstrated that the
uniform distribution of highly dispersed CuNi particles and surface basicity were crucial for the enhanced catalytic performances [93].

3.3. Hydrogenation of Levulinic Acid

Levulinic acid (LA) is an important biomass platform chemical which can be produced by the acid-catalyzed hydrolysis of lignocellulose [94]. One of the most important downstream chemicals from LA is gamma-valerolactone (GVL), a stable and nontoxic lactone that has broad applications and could be further converted into valuable compounds [95]. For example, GVL could be transformed into 1,4-pentanediol (1,4-PDO) by a hydrogeneative ring-opening reaction, and 1,4-PDO could go through an intermolecular dehydration to form 2-methyltetrahydrofuran (2MTHF) [26,96]. Two mechanisms for LA hydrogenation into GVL were proposed based on literature research. In the first mechanism [97], LA was initially hydrogenated to form 4-hydroxypentanoic acid on metallic sites, followed by the lactonization to yield GVL at acid or basic sites. The second mechanism [98] started with an isomerization of LA to the enol form, followed by the lactonization to produce angelica lactone, and ends with hydrogenation to produce GVL. Some studies stated that the hydrogenation was the rate-determining step, while the lactonization reaction proceeded rapidly [99]. High hydrogen pressure (0.5–3 MPa) is usually necessary for high conversion of LA, so that the sufficient amount of dissolved hydrogen in solvent enables LA hydrogenation [100]. Many catalysts based on LDHs or MMOs were reported to be efficient for converting LA to GVL with high yields. Based on the published research, it seems that MMO catalysts based on Cu, Ni or Co showed comparably high selectivity toward GVL from LA. For example, CuAl-MMO [82], NiAl-MMO [101,102] and CoAl-MMO [103] catalysts were synthesized by different research groups and all showed comparably high selectivity (>85%) toward GVL. Analogously, CoMoAl [104], NiMgAl [101] and CuMgAl [105] catalysts synthesized by calcination of LDH precursors also showed dominant GVL production. Noble metal supported on LDH or MMO materials were also efficient for LA hydrogenation. Notably, MMOs-supported noble metal catalysts with a much lower reaction temperature (40–80 °C) were reported recently, compared to a reaction temperature of 140–260 °C used by most non-noble metal catalysts. Pt supported on MgAl-MMO demonstrated a nearly complete LA conversion into GVL at 40 °C and 50 bar [106], while Ru supported on MgAl-MMO or MgLa-MMO also showed a GVL selectivity higher than 99% at 80 °C and 5 bar [107].

Although high selectivity toward GVL from LA was achieved by many catalysts, a major concern in LA hydrogenation catalysts that should be dealt with is catalyst deactivation. Several mechanisms were proposed, including particle sintering, coke deposition, or metal leaching [108,109]. Because LA conversion into GVL produces water in lactonization, it is probable that MgO and Al₂O₃ may undergo phase transition in the hydrothermal environment, with brucite MgO or γ-Al₂O₃ transformed into periclase or boehmite, respectively, leading to undesired changes in pore structure and surface property [110–112]. Deactivation of LDH or MMO catalysts in LA hydrogenation was also observed and studied for its mechanisms. Hu’s research reported Ni leaching in NiMgAl-MMO catalysts in LA hydrogenation, mainly due to the detrimental effect of water to Mg or Al leading to structure collapse [101]. Hussain et al. observed significant decreases of LA conversion in a time-on-stream study of MgAl-MMO. After characterization on spent catalysts, it is confirmed that in addition to the coke deposition on catalyst surface, the MgAl-MMO catalyst adsorbed water to regain the layered structure due to the memory effect, though this deactivation was largely reversible by calcination at 450 °C in air [113]. Considering the structure destruction during the reaction, metal particle sintering becomes inevitable, as observed with Cu/MgAl-MMO catalysts that Cu particles sized increased from 2 to 5.3 nm after LA hydrogenation at 260 °C [114]. Therefore, the stability of LDH or MMO catalysts for LA hydrogenation is an unavoidable issue before the commercialization of the catalytic process.
3.4. Catalytic Hydrogenation of Monosaccharides

The conversion of biomass feedstocks to fuels or chemicals is an important approach to generate renewable energy resources. Lignocellulose depolymerization or fermentation could produce monosaccharides or sugar alcohols, which could be further transformed to various types of chemicals [115]. For example, cellulose or hemicellulose could be hydrolyzed into C$_5$ and C$_6$ sugar alcohols such as glucose, xylose, and arabinose. These sugar alcohols were listed by the US Department of Energy among the top 12 sugar-derived building block chemicals, which are stable and versatile to upgrade to commodity chemicals [116]. Yamaguchi et al. prepared hydrotalcite supported nickel phosphide with high activity and stability for the hydrogenation of glucose [117], xylose [118], and maltose [119] under mild condition. The authors indicated that the hydrotalcite support activated the carbonyl group of the saccharide molecules and donated electrons to Ni, while molecular hydrogen was activated by unsaturated Ni sites from Ni$_2$P. Another study on sugar hydrogenation using a physical mixture of PtSn/Al$_2$O$_3$ and MgAl-MMO proposed that the addition of MgAl-MMO could provide an alkaline environment in the reaction system to facilitate ring opening reactions of sugar molecules [120], which is a key step for carbonyl hydrogenation [121]. While supported LDH catalysts showed synergistic effects between support and active metal, LDH-derived MMOs were also active in sugar hydrogenation. Wu et al. have shown that reduced CuNiAl and CuNiAlM (M = Mg, Co, Cr and Fe) catalysts were effective for the hydrogenation of glucose and fructose into sorbitol and mannitol [122,123]. Pérez-Ramírez et al. also reported HT-derived CuNiAl catalysts could catalyze transfer hydrogenation of these sugars to the corresponding polyols [124]. In both research, characterization results indicated that metallic Cu and Ni were formed in the reduced catalysts, which could act as active sites for hydrogenation.

4. Hydrogenation of Carbon-Carbon Unsaturated Bonds

4.1. Partial Hydrogenation of Alkynes

The production of ethylene in oil refinery is often accompanied with acetylene as the byproduct. Because acetylene could poison Ziegler–Natta catalysts and significantly deteriorate the quality of polyethylene, a partial hydrogenation process is needed to eliminate ethylene and not overhydrogenate ethylene to ethane [125]. Acetylene hydrogenation is considered to be a structure-sensitive reaction in which catalyst composition plays a vital role in product selectivity [126]. Ma et al. obtained a Pd/MgAl-LDHs/Al$_2$O$_3$ catalyst by in situ synthesis of Pd/MgAl-LDHs on spherical Al$_2$O$_3$ surface, and another Pd/MgO-Al$_2$O$_3$ catalyst by calcination and reduction of synthesized Pd/MgAl-LDHs/Al$_2$O$_3$ [127]. The results indicated that both catalysts had larger surface area, lower surface acidity, uniform Pd particle size, and strong metal-support interaction, leading to higher catalytic activity and selectivity than impregnated Pd/Al$_2$O$_3$ [128]. The high dispersion of Pd on the HT surface is presumably originated from the acidic sites on the support surface which weakened the electron density of Pd, favored the formation of low-coordinated Pd sites and enhanced dispersion [129,130]. Because Ag could effectively inhibit the formation of coke and improve the activity and stability of the catalyst [125], PdAg bimetallic catalysts supported on LDH-derived MMOs were also studied for acetylene hydrogenation. Research on PdAg supported on LDH-derived ZnO-Al$_2$O$_3$ found that compared with Al$_2$O$_3$, ZnO-Al$_2$O$_3$-MMO could significantly inhibit the overhydrogenation of acetylene and decrease oligomer formation [131]. Research on PdAg supported by NiTi-MMOs for acetylene hydrogenation found that a large number of Ti$^{5+}$ defective sites existed on the NiTi support, acting as active centers to activate hydrogen and increasing the electron density Pd, thus promoting the desorption of ethylene and improving reaction selectivity [132]. When switching NiTi to MgTi as the PdAg support, the electronic effect between Ti$^{5+}$ and Pd was still present, combined with the acid–base property of the MgTi support, contributing to nearly 100% conversion and 83.8% selectivity to ethylene at 70 °C [133].

In addition to being used as a catalyst support, LDH materials after calcination and reduction can also be used directly as acetylene hydrogenation catalysts. Rives et al. [134,135]
prepared a series of NiZnAlCr hydrotalcite-like catalysts for the selective hydrogenation of acetylene. The introduction of Cr inhibited the formation of coke, while the addition of Zn enhanced the metal-support interaction and improved the dispersion of Ni. Liu et al. [136] prepared CuNiMgAl nanoalloy catalysts for acetylene partial hydrogenation using LDHs as the precursor. High metal dispersion and structural homogeneity of NiCu nanocrystals were observed, and this LDH-derived CuNiMgAl showed better selectivity, anti-coking ability and stability than the CuNi/MgAl-HT catalysts prepared by the impregnation method. The addition of iron in Cu-based MMOs could further improve selectivity and reduce oligomerization, because iron served as a structural promoter to disperse active metals [137]. Moreover, in another research investigating Cu/Fe\textsubscript{y}MgO\textsubscript{x}-type catalysts derived from LDHs, iron is proposed to facilitate the formation of bifunctional interfacial active site Cu\textsuperscript{δ−}Fe\textsubscript{0.16}\textsuperscript{δ+}MgO\textsubscript{x}, which is effective in activating acetylene and hydrogen and promoting the desorption of ethylene [138].

Similar to acetylene partial hydrogenation, the partial hydrogenation of phenylacetylene to styrene for the elimination of phenylacetylene is an important pretreatment step in styrene polymerization process. Several LDH-based catalysts have shown excellent performance in this reaction. Pd/HT catalysts were synthesized for the phenylacetylene hydrogenation, and it is proposed that the layered structure of HT may impose steric restriction on reactants, making Pd/HT more stereoselective in the product than the conventional supported Pd catalysts [139]. Duan’s group presented a novel series of nickel phosphide catalysts from LDH precursors with high selectivity toward styrene. It is claimed that phosphorus incorporation increased the Ni–Ni bond length and decreased Ni electron density, leading to the desorption of styrene so that further hydrogenation was avoided [140].

4.2. Hydrogenation of Aromatic Ring

Aromatic compounds in transportation fuels are responsible for reduced cetane number and increased particulate emission, as well as potential health hazard accompanied by the emission [141–143]. Strict legislative restrictions on aromatic content all over the world encourage research efforts on aromatic rings hydrogenation as an effective strategy to eliminate aromatic compounds in fuels [144]. The urgent need for lignin upgrade in recent years also inspires research on efficient aromatic hydrogenation [145]. Bai et al. investigated the hydrogenation of phenol over a series of Ni-based MMO catalysts and over 90% selectivity of cyclohexanol was obtained at 110–150 °C [146–148]. It is pointed out that the strong metal–support interaction of MMO catalysts induced high metal dispersion and prevented agglomeration or sintering. Strong metal–support interaction was also important in aromatic hydrogenation over Pd supported by MMOs [149]. Highly dispersed Pd particles were formed on CoCeAl-MMO surface with a particle size smaller than 4 nm, providing plentiful active sites for H\textsubscript{2} dissociative adsorption and ring hydrogenation. The addition of Ce generated abundant oxygen vacancies, which activated phenol by forming phenoxy and accelerated the hydrogenation reaction. Reaction temperature and pressure also significantly affected hydrogenation activity, as relatively high temperature and high pressure were more preferable for benzyl ring hydrogenation products [150,151].

5. Hydrogenolysis of Oxygenated Compounds

5.1. Hydrogenolysis of Esters

Ester hydrogenation and hydrogenolysis are important reactions with particular interest in biomass upgrade. Esters, including fatty acid esters, lactones, and levulinate esters, could be either directly extracted or chemically converted from biomass feedstock. Because esters themselves have limited uses in the chemical industry, hydrodeoxygenation or hydrogenolysis is usually necessary to convert esters into valuable commodities such as alcohols or alkanes. Triacylglycerols from non-edible plant oils or animal fats could be converted into fatty acid esters by transesterification reactions. Due to their high oxygen content and low heating value as fuels [152], researchers attempted to upgrade fatty acid esters to diesel-range hydrocarbon (C\textsubscript{15}–C\textsubscript{18}) by hydrodeoxygenation. Cao et al. utilized
LDH-derived NiCuAl catalysts for the hydrogenolysis of soybean and waste cooking oils, and more than 80% yield for diesel-range hydrocarbons was obtained at 260 °C and 3 MPa of H₂ [153]. Lewis acid sites (Al³⁺) was thought to be active sites for decarbonylation by interacting with the oxygen atom and cleaving acyl C-O bond. Another study [154] using NiGaMgAl to catalyze the hydrodeoxygenation of methyl laurate (C₁₁H₂₃COOCH₃) also present a 99% yield toward C₁₁+C₁₂ hydrocarbons at 400 °C and 3 MPa. The promotional effect of Ga to Ni suppressed the C-C bond hydrogenolysis, accounting for the high hydrocarbon yield.

Converting esters to diols which could be used as monomers for polyester materials is another attractive tactic with practical needs. Ethylene glycol, an important monomer in the polymer industry, could be produced by an indirect pathway from syngas via dimethyl oxalate. Wei’s group prepared CuZrMgAl-MMO catalysts for dimethyl oxalate hydrogenation and obtained 99.5% yield of ethylene glycol under 180 °C and 2 MPa [155]. According to in situ characterization techniques, the authors attributed the high reactivity to the Cu-O-Zr metal-support interfacial sites, where carbon–oxygen bonds were activated at Zr-related oxygen vacancies and hydrogenated at adjacent Cu sites. Li’s group demonstrated that 1,4-pentanediol could be produced from bio-derived ethyl levulinate by hydrodeoxygenation over CuCoAl-MMO catalysts, where Lewis acid sites such as electrophile Cu⁺ or electron-deficient CoOₓ were considered to be crucial for carbonyl adsorption and activation [25].

5.2. Hydrodeoxygenation of Lignin Derivatives

Lignin is a cross-linked three-dimensional amorphous polymer composed of aromatic units, accounting for 10–35 wt.% in lignocellulosic material [145,156]. The rapid development of the biomass industry cogenerated more than 100 million tons of lignin annually [157]. Various aromatic chemicals could be produced by lignin valorization, yet only a small portion of lignin was actually utilized for chemical production. Lignin depolymerization is a complex and challenging task, with the pivotal issue being the cleavage of C-C bond and C-O bond. Hydrogenolysis is an effective way to break C-O linkage, and various LDH-derived catalysts were demonstrated to be highly reactive for lignin hydrogenolysis due to its high metal dispersion and adjustable acid–base properties. Flower-like Ni₂P-Al₂O₃ catalysts were obtained by reduction of red phosphorus with NiAl-LDH precursors at 500 °C by Li’s group [158]. Ni₂P-Al₂O₃ showed high alkane selectivity in hydrogenolysis of poplar lignin oil. The high reactivity was attributed to the high exposure of cantilevered conical Ni sites with strong C-O bond break capability and plentiful acid sites to activate the substrate and H₂, as shown in Figure 3. Other research stated that acid sites, especially Lewis acid sites, facilitated deoxygenation of lignin derivatives by adsorbing the electron-rich phenyl ring or oxygen atoms, promoting ring hydrogenation or deoxygenation of methoxy or carbonyl [159,160]. Oxygen vacancies are also important for constructing high reactivity LDH-derived catalysts for lignin hydrodeoxygenation. Reducible oxide such as CoOₓ [161] or CeOₓ [162] in MMO catalysts could form surface oxygen vacancies to adsorb or activate oxygen-containing functional groups, greatly benefiting lignin deoxygenation reaction.

Among the common linkages that exist in lignin, the most recurring linkage is the β-O-4 linkage, constituting 50% of all linkages [145]. Therefore, model compounds with β-O-4 linkages were often synthesized to investigate catalyst reactivity in lignin depolymerization [163–165], Wang et al. prepared a NiMgAl-C composite catalyst using lignosulfonate as the carbon precursor to impose electronic modifications to Ni [164]. The synthesized catalysts showed promising results in terms of conversion and selectivity in hydrodeoxygenation of model compounds and real lignin. The lamellar structure of LDHs could also be utilized to improve catalytic performance. Beckham and colleagues presented an inspiring study, where HT-derived catalysts with intercalated nitrates produced phenol monomers 2–3 times more than catalysts without nitrates in hydrodeoxygenation of model compounds [166]. It is proposed that intercalated nitrates not only increased accessibility to
basic active sites but also directly participated in the hydrogenolysis process, as the activity could be recovered after the nitrate reservoir was depleted and replenished.

![Reaction mechanism of guaiacol hydrodeoxygenation over NiP catalysts](image)

**Figure 3.** Reaction mechanism of guaiacol hydrodeoxygenation over NiP catalysts [158]. (Copyright 2022, American Chemical Society).

### 5.3. Hydrodeoxygenation of Furfural

C₅ polyols are important types of chemicals with huge demands in the polymer industry. Such polyols are traditionally produced from petroleum resources. Recently, researchers have developed the catalytic transformation of furfural to various polyols via hydrodeoxygenation. For furfural conversion to pentanediols (PDO), although early in 1930s Adkins and Connor presented Cu-Cr catalysts to convert furfuryl alcohol to pentanediols [167], MMO catalysts for catalytic conversion of furfurals to pentanediols were extensively studied in the last decade. Initially, Pt supported on CoAl-MMO [168] or MgAl-HT [169] synthesized by two separate research groups showed high reactivity in furfural hydrogenolysis toward 1,5-pentanediol and 1,2-pentanediol, respectively, at 110–140 °C and 1.5–3 MPa. The role of these supports is related to the basicity which might change adsorption preference leading to ring-opening reactions. Subsequently, Huang’s group noticed that CuMgAl-MMO [170] or CuAl-MMO [171] alone could also catalyze the formation of pentanediols at a relatively high pressure of 6 MPa. The distinct difference in reactivity and selectivity between conventional impregnated Cu/Al₂O₃ catalyst and coprecipitated CuAl-MMO catalyst demonstrated the advantage of the much smaller Cu particle size of the MMO catalyst (1.9 nm) compared to the impregnated catalyst (16.7 nm) [171]. Another advantage of MMO catalysts is the partially reduced metal species on the surface, as several studies presented positive correlations between partially reduced metal species and pentanediol selectivity [24,172,173]. Based on published literature, the mechanism of furfural conversion to pentanediols could be summarized, as shown in Figure 4. Furfural is first hydrogenated to produce furfuryl alcohol, and then furfuryl alcohol will go through C-O bond scission to afford 1,2-pentanediol or 1,5-pentanediol depending on the catalyst property. Acid–base active sites [24,174–177] or partially reduced metal species [172,173] are proposed to influence the adsorption configuration of furfuryl alcohol, resulting in the activation of different C-O bond and different pentanediol products.
5.4. 5-Hydroxymethylfurfural Hydrodeoxygenation

Cellulose could be transformed into 5-hydroxymethylfurfural (HMF), a platform chemical with various downstream products [77]. Many important compounds can be produced from HMF. For example, 2,5-dimethylfuran (DMF) by HMF hydrodeoxygenation could be used as fuel additive due to its high-octane number and energy density [178,179]. HMF could also be converted to 2,5-furandimethanol (FDM) [180,181] via hydrogenation of carbonyl group, to 2,5-tetrahydrofuran dimethanol (THFDM) [180,182] via furan ring hydrogenation of FDM, or to 2,5-dimethyltetrahydrofuran (DMTHF) [183–185] via furan ring hydrogenation of DMF. FDM and THFDM could be used as building blocks in polymer synthesis, and DMTHF is a potential fuel additive [77]. As shown in Table 3, the selectivity toward DMF, FDM, or DMTHF could be controlled by choosing adequate reaction conditions or metal catalysts [180,184].

Table 3. Catalytic performances of LDH-derived catalysts for 5-hydroxymethylfurfural hydrodeoxygenation 1.

| Catalyst         | Reaction Condition | Conversion (%) | Major Product Selectivity (%) | Ref.   |
|------------------|--------------------|----------------|------------------------------|--------|
| NiAl-MMO         | 80 °C; 2 MPa H₂    | 96.0           | THFDM, 74                    | [180]  |
| Ru/MgAl-HT       | 220 °C; 1 MPa H₂   | 100.0          | DME, 58                      | [181]  |
| NiAl-MMO         | 180 °C; 1.2 MPa H₂ | 100.0          | DMTHE, 97.4                  | [182]  |
| Ni-Cu/HT         | 90 °C; 1 MPa H₂    | 99             | DME, 67                      | [185]  |
| CoZnAl-MMO       | 130 °C; 0.7 MPa H₂ | >99.9          | DME, 74.2                    | [186]  |
| CuCoNiAl-MMO     | 180 °C; 1 MPa H₂   | 99.8           | DME, 95.3                    | [187]  |
| CuZnAl-MMO       | 180 °C; 1.2 MPa H₂ | 100            | DME, 90.1                    | [188]  |
| Co-N-C/NiAl-MMO  | 170 °C; 1.5 MPa H₂ | 99.9           | DME, 100                     | [189]  |
| NiZnAl           | 100 °C; 1.5 MPa H₂ | 100            | FDM, 98.2                    | [190]  |
| Cu@Co/CoAlOₓ     | 180 °C; 1.2 MPa H₂ | 100            | DHTMF, 83.6                  | [191]  |
| CuCoCe-MMO       | 210 °C; 1.5 MPa H₂ | 100            | DME, 96.5                    | [192]  |
| NiCoAl-MMO       | 120 °C; 4 MPa H₂   | 100            | 1,2,6-HTO, 64.5              | [193]  |
| Cu₁₃Mg₁₅Al       | 150 °C; 6 MPa H₂   | 100            | 1,2-HDO, 40                  | [194]  |

1. THFDM, 2,5-tetrahydrofuran dimethanol; FDM, 2,5-furandimethanol; DMF, 2,5-dimethylfuran; DMTHE, 2,5-dimethyltetrahydrofuran; 1,2,6-HTO, 1,2,6-hexanetriol; 1,2-HDO, 1,2-hexanediol.

Figure 4. Reaction mechanism of furfural hydrogenolysis over CuMgAl–MMO catalysts [170]. (Copyright 2022, Royal Society of Chemistry).
The transformation of HMF to afford DMF could be accomplished by various metal catalysts. CuAl- [80], NiAl- [180,182], and CuNiAl-MMO [195] catalysts with high metal dispersion showed excellent performance for the conversion of HMF to DMF, with DMF yields higher than 90%. In contrast, CoAl showed relatively lower DMF selectivity [186,187], likely due to its lower hydrogenation reactivity [196]. The conversion of HMF into DMF requires multifunctional catalysts with dual active sites catalyzing C=O hydrogenation and C-O hydrogenolysis. Wang et al. [188] studied the HMF hydrodeoxygenation over CuZnAl-MMO catalysts, claiming that Cu⁺ species acted as C-O cleavage sites and Cu⁰ species acted as C=O adsorption sites. The synergy between Cu⁺ and Cu⁰ species promoted the selective transformation of multi-functional groups of HMF molecules, thus improving catalyst reactivity and selectivity. However, these active sites might be susceptible to deactivation caused by changes in the metal chemical state [189]. This issue could be tackled by the incorporation of Zn generating strong Cu-O-Zn interaction by charge compensation, stabilizing active Cu⁺ species, influencing adsorption configuration and preventing Cu agglomeration [186,190]. In another study on LDH-derived CuCoAl catalysts, Wang et al. pointed out that interfacial sites played key roles in the formation of different reaction intermediates in the catalytic hydrodeoxygenation of HMF [191]. C=O bond in the reactant was hydrogenated by the oxygen vacancies from CoOx site to form hydroxy groups, while C=C bond was hydrogenated by Cu-Co interface positions. A similar interfacial effect is observed in reduced CuCoCe [192] and CoMgFe [197] catalysts as well, where oxygen vacancies formed on CuCeOx or CoFeOx interface activated the C-O bond and promoted C-O cleavage.

The hydrogenolysis of HMF to hexanediols or hexanetriols over LDH-derived catalysts were also probed by researchers. NiCoAl-MMO [193] and Pt supported on MgAl-MMO [198] were effective in converting HMF to 1,2,6-hexanetriol at 120–160 °C and 3–4 MPa. A 64.5% yield was obtained with a 0.5Ni2.5CoAl catalyst at 120 °C and 4 MPa. 1,2-Hexanediol could be produced over CuMgAl-MMO catalysts at 150 °C and 4–6 MPa [194]. The structure–activity relation could be inferred by two correlations reported in Hu’s research [194], as shown in Figure 5: a positive correlation between basic sites and catalyst turnover frequency (TOF), and a negative correlation between Cu particle size and catalyst TOF. These observations hinted at the crucial role of Cu particle size and basicity, which controlled the catalysts reactivity and selectivity in HMF hydrodeoxygenation, analogous to furfural hydrogenolysis. Conceivably, HMF or its hydrodeoxygenated product 5-methylfurfuryl alcohol were adsorbed on active sites in a tilted mode with C-O bonds interacting with active sites, leading to the C-O bond cleavage to form polyols [193,199].

Figure 5. Influence of the molar ratio of Cu/Mg on the (a) distribution of basic sites and (b) Cu particle size. (c) Correlation between the Cu particle size and TOF value and (d) basic sites and TOF value [194]. (Copyright 2022, American Chemical Society).
6. CO₂ Hydrogenation

The chemical fixation of CO₂ into value-added products is a pivotal process in CO₂ utilization to reduce CO₂ emission, decrease fossil fuel usage, and alleviate the greenhouse effect. Numerous research on catalytic conversion of CO₂ were conducted in recent years, and exciting progress was made for the hydrogenation of CO₂ into various chemicals including methane, methanol, carbonates, carboxylic acid, etc. [200, 201]. Compared to regular hydrogenation catalysts, unique advantages in adsorption capacity of LDH-derived catalysts for CO₂ hydrogenation are recognized to further promote catalyst performance through adsorption enhancement [202]. Because a number of reviews were published in the past few years regarding the broad CO₂ hydrogenation field [9, 201, 203, 204] or specific CO₂ hydrogenation reaction [205], the goal of this review is to analyze the applications of LDH or MMO catalysts in CO₂ hydrogenation, focusing on CO₂ thermocatalytic conversion into methanol and methane.

6.1. CO₂ Conversion to Methanol

The CO₂ hydrogenation to methanol is a promising strategy for CO₂ utilization, as methanol is a key feedstock that can be industrially converted to a wide range of chemicals such as light olefins and gasoline [206, 207]. For example, the methanol-to-olefin (MTO) process by Dalian Institute of Chemical Physics, China, has been commercialized with an annual production of 7.16 Mt per year by the end of 2018 [208]. Methanol could be produced from syngas (CO₂/CO/H₂) using an industrial Cu/ZnO/Al₂O₃ catalyst synthesized from zincian malachite, (Cu,Zn)₂(OH)₂CO₃, by the coprecipitation of metal nitrates (Cu, Zn and Al) [209]. It has been conclusively demonstrated that the superior reactivity of Cu/ZnO/Al₂O₃ stemmed from the structure defects of the Cu surface and the promotional effect of Zn to Cu [210]. Owing to the advantages of the MMO catalysts mentioned above, research efforts were devoted to preparing CuZnAl-based catalysts from LDH precursors aiming to obtain catalysts more reactive than the industrial Cu/ZnO/Al₂O₃ catalyst. Behrens and coworkers presented CuZnAl-MMO catalysts through the coprecipitation of nitrate solution at 25 ºC and a constant pH value of 8 ± 0.7. The synthesized CuZnAl was intrinsically more active in CO₂ hydrogenation than the industrial CuZnAl catalyst when normalized by Cu surface area, due to the highly dispersed Cu particles (7 nm) and strong interfacial interaction between embedded Cu particles and ZnAl₂O₄ matrix [211].

To further improve the catalyst reactivity of CuZnAl catalysts, the addition of fourth metal elements was explored in CO₂ hydrogenation to methanol. A wide range of metals have been added to CuZnAl by coprecipitation of metal precursors, with different metals exhibiting different modification effects. The addition of zirconium to CuZnAl could increase the basicity, strengthen the adsorption of CO₂ and promote the formation of methanol [212, 213]. Similar effects were observed for Mn, La, Ce and Y, as a series of CuZnAlX (X = Mn, La, Ce, Zr and Y) catalysts were synthesized in a study from Sun’s group, and modified CuZnAlX catalysts showed a higher portion of strong basic sites, CO₂ conversion and methanol selectivity than the unmodified CuZnAl [214]. Yttrium modified CuZnAl possessed a higher surface area and more dispersed Cu particles, resulting in a higher methanol yield per gram of catalysts than the unmodified CuZnAl [215]. The addition of indium to CuZnAl increased the methanol selectivity by inhibiting CO formation, yet decreased TOF of methanol formation [216].

Modifications on synthesis procedures were also attempted to induce structural transformation of the CuZnAl-based MMO catalysts. Different alumina sources [217, 218], Zn precursors [219, 220], phase of precipitated precursors [221], atomic ratios [222–224], sequence of precursor addition [225, 226], calcination temperature [227], precipitation pH [228] and precipitation agents [229] in synthesis procedures of MMO catalysts commonly resulted in diverse catalytic activity, selectivity or stability for CO₂ hydrogenation. These phenomena were often attributed to the difference in acid–base properties, exposed Cu surface area, particle size, phase composition, etc. However, due to the structural complexity of active sites in methanol synthesis [210, 230], it is not easy to acquire a direct correlation
between catalyst physiochemical properties and catalytic performance. More in-depth research on catalyst structure, especially in situ characterizations, is essential to unveil the structure–activity relationship for CuZnAl catalysts.

For the industrial Cu/ZnO/Al2O3 catalyst, the role of alumina is considered to be a structural promoter that prevents Cu particles from aggregation [231]. Kühl et al. conducted experiments to partially replace Al with trivalent cations Cr and Ga [232]. The substitution of Al by Cr reduced the interaction between Cu and oxide matrix, causing Cu particle growth during CO2 hydrogenation. In contrast, the substitution of Al by Ga turns out to be rather effective in improving catalyst activity, and this improvement was ascribed to the fact that Ga addition could stabilize Cu phase and benefit the catalytic activity of Cu particles. Other research presented consistent conclusions. CuZnGa-MMO catalysts showed higher Cu surface area and dispersion than CuZnAl using the aqueous miscible organic solvent treatment (AMOST) method, which involves an additional step of treating LDHs precipitates with an organic solvent [233,234].

6.2. CO2 Conversion to Methane

The conversion of CO2 to methane, namely CO2 methanation, is also known as the Sabatier reaction first reported in 1897. In the current age of pursuing carbon neutrality, Sabatier reaction is experiencing its renaissance, as CO2 methanation becomes a convenient strategy to convert CO2 to a widely used energy resource in a sustainable manner. Two types of reaction mechanisms for CO2 methanation were proposed. The first mechanism presumes the direct methanation of CO2 to methane via formate, carbonate or bicarbonate, while the second mechanism requires the conversion of CO2 to CO before CO is converted to CH4 [235–237]. The actual mechanism may vary depending on the catalyst type, reaction temperature, etc. As CuZn-based catalysts constitute the majority of catalysts for methanol synthesis, Ni-based catalysts represent the most important category of CO2 methanation catalysts [236]. Another important factor for the catalyst of methane formation is the amount and strength of basic sites [238,239]. In this respect, MMO catalysts derived from MgAl-LDHs were studied comprehensively due to its advantageous capability of combining tunable basic sites with dispersed metal particles.

The strong dependence of CO2 conversion to catalyst basicity, especially medium strength basic sites, was reported by many researchers, as indicated by the positive correlation between medium strength basic sites and CO2 conversion or TOF [240–242]. Basic sites, such as OH− and O2−, could adsorb and activate CO2 to form carbonate or bicarbonate species [243,244]. It is noticed that increased basicity was related to the increased amount of bicarbonate-like species for NiMgAl-MMO catalysts, while bicarbonate is the main CO2 adsorption product and an important intermediate for methane formation at low temperatures [245]. It is also speculated that higher basicity promotes the hydrogenation rate of surface-adsorbed carbonates [246]. Moreover, medium strength basic sites promote monodentate carbonates formation, as monodentate is more readily hydrogenated to form CH4 than bidentate carbonates [238,245]. The introduction of certain metal promoters is a facile way to manipulate MMO catalyst basicity. Metals including V [247], Y [240], Fe [246,248], Mn [242], Zr [249], Ce [250], La [251–253] and Cu [241] have been demonstrated to result in increased concentration of basic sites. Apart from changes in basicity, metal promoters could also adjust the metal–support interaction and increase Ni dispersion. Ho and colleagues [253] claimed that CO2 interacted more strongly with NiLaAl than NiAl due to higher basicity and more dispersed Ni particles, leading to a much higher low-temperature activity in CO2 methanation of NiLaAl. In situ characterization studies suggested that both dissociative activation (forming CO) and associative activation (forming bicarbonate and carbonate) of CO2 were simultaneously observed for NiLaAl, while for NiAl, CO2 was successively activated by the dissociative pathway and then associative.

Oxygen vacancies are also proposed as a crucial factor for CO2 methanation, as shown in Figure 6 [250]. Oxygen vacancies could act as Lewis acid sites to interact with electron-rich oxygen atoms from CO2 for CO2 chemisorption and activation [254]. DFT calculations...
confirmed that oxygen vacancies on simulated the NiCeAl surface contributed to the lower CO$_2$ adsorption energy than the NiAl surface [255]. Oxygen vacancies also facilitate the formation of active oxygen species, which could react with CO$_2$ to produce monodentate or bidentate carbonates, accelerating the methanation reaction [249,250]. He et al. confirmed that NiZrAl ternary MMO catalysts showed more oxygen vacancies and basic sites than NiAl binary counterpart.

**Figure 6.** The role of oxygen vacancies in CO$_2$ methanation via (a) formate pathway and (b) CO pathway [250]. (Copyright 2022, Elsevier).

**7. Hydrogenation in C-C Coupling Processes**

In recent years, the demand for bio-derived fuels or chemicals stimulates extensive interests in converting biomass feedstock into various types of renewable fuels. Regarding the fact that most biomass platform chemicals consist of oxygen-containing functional groups such as hydroxy or carbonyl groups, C-C coupling or C-C bond formation via aldol condensation, dehydration/hydrodeoxygenation, and hydrogenation are effective strategies to produce downstream chemicals, especially in the synthesis of long chain hydrocarbons such as jet fuel in the range of C$_8$–C$_{16}$. Regarding the advantages of LDH-derived MMOs simultaneously possessing metallic, acidic and basic sites, they become ideal candidates either as catalyst supports or as the catalyst themselves to achieve the cascade hydroconversion of C-C coupling for biomass upgrade. In some cases [256–262], C-C coupling to produce chemicals was accomplished by a two-stage process, with the first stage aimed at aldol condensation and dehydration using HT or MMO catalysts, and the second stage aimed at hydrogenation using non-HT and non-MMO catalysts. These studies, which did not use HT or MMO as hydrogenation catalysts, will not be discussed in this review. The following section will examine literature work on HT- or MMO-catalyzed hydrogenation in the C-C coupling processes.

Acetone, an inexpensive byproduct in lignocellulose fermentation, could undergo aldol condensation with itself or other chemicals to produce products with longer chains. The transformation of acetone to methyl isobutyl ketone (MBK), an important chemical with extensive applications, is a typical example of the multifunctionality of LDH-derived materials. MBK could be converted from acetone via three consecutive steps: aldol condensation catalyzed by bases, dehydration catalyzed by acids, and hydrogenation catalyzed by metals [263]. Regarding the advantages of LDH-derived MMOs, they are catalytically active as supports (Pd on ZrCr [263], MgAl [264,265] and CoAl [266]) or catalysts themselves (NiMgAl [267] and CuAl [268]) to achieve the one-pot conversion of acetone to MBK. Because the sequence of the three steps are crucial for MBK production from acetone, the concentration and the strength of acidic, basic, and metallic sites are expected to be balanced in order to minimize side products [264]. In order to obtain the optimal catalysts, catalysts with different concentrations of acid or base sites were prepared and tested. According to experimental results, it is generally believed that excessively
high acidity or basicity would lead to over-condensation products and reduce MIBK yields [264,267].

Cyclopentanone (CPO, C₅H₈O), a bio-derived chemical produced by aqueous phase furfural rearrangement, could be transformed to jet-fuel range cycloalkanes by C-C coupling reactions. Cai and coworkers synthesized a bifunctional Ni/MgAl/active carbon catalyst using a MgAl-HT precursor in an integrated C-C coupling/hydrogenation process for the conversion of CPO with over 80% yield toward C₁₀ or C₁₅ alkanes [269,270]. The authors ascribed the remarkable performance to the enhanced strength of the basic sites of MgAl-MMO, promoting trimerization of CPO more effectively than conventional oxide support such as MgO, Al₂O₃, NaY or TiO₂, as well as the strong hydrodeoxygenation activity of Ni than Fe, Co or Cu. Another research combining MgAl-MMO with Pd in CPO trimerization also obtained high trimer yield [271]. In addition to CPO, C-C coupling of other ketones were also attempted to produce elongated alkanes using LDH or MMO catalysts. Sheng et al. performed MIBK self-condensation over Pd-modified MgAl-HT to produce dodecanol, with MgAl-HT support acting as a base catalyst for self-aldol condensation [272].

Alcohols could also be used for C-C coupling after dehydrogenation to form aldehydes or ketones, followed by aldol condensation, dehydration and hydrogenation. For these processes, metallic sites on MMO catalysts could catalyze the dehydrogenation of the initial alcohol and the hydrogenation of the unsaturated ketones, while acid–base sites catalyze condensation and dehydration reaction. A few alcohols were studied for their C-C coupling applications, including 2-hexanol [273], propanol [274], octanol [275] and ethanol [276,277]. It is commonly believed in these studies that numbers of metal, acid, and base sites should be delicately designed to prevent side reactions and by-products.

8. Hydrogenation of Nitrites and Nitriles

8.1. Hydrogenation of Organic Nitrites

The selective hydrogenation of aromatic nitro compounds to produce anilines is important in synthetic chemistry. The major challenges in this process are to selectively reduce nitro groups and not affect other functional groups. Various types of noble or non-noble metal catalysts were developed to achieve this goal [278]. For catalysts with noble metals such as Pd or Pt, key factors that determine their catalytic performance are metal dispersion and the resistance to particle agglomeration. LDHs or MMOs, which exhibit hierarchical structure able to immobilize active species, stand out as desirable catalyst support. Wang et al. showed that Pd supported on MgAl-LDHs demonstrated higher catalytic activity and selectivity than Pd/SiO₂ or Pd/C in substituted nitrobenzene hydrogenation, likely due to the confining effect of metallic Pd in layers of LDHs [279]. Similar conclusions were reached in another study studying Pd supported on MgAl-MMOs, MgO and γ-Al₂O₃ in nitrobenzene hydrogenation [48]. Results showed that Pd/MMOs were superior to the latter two in terms of turnover frequency and stability, and authors ascribed the superiority to the high dispersion of Pd particles on MMOs and the anchoring effect of MMOs to prevent Pd aggregation. Non-noble MMO catalysts such as CuZnAl also showed high activity in the hydrogenation of nitro groups [280]. The activity of CuZnAl was attributed to the Cu-ZnOₓ active sites, similar to the one used in CO₂ conversion, as mentioned above.

A more challenging case in aromatic nitro compound hydrogenation is the selective hydrogenation of nitro groups when other readily reducible groups are presented, as in the case of nitrostyrene hydrogenation to yield vinylaniline. Corma and Serna have reported a breakthrough by using Au/TiO₂ and Au/Fe₂O₃ to hydrogenate 3-nitrostyrene to selectively produce 3-vinylaniline [281]. Recently, a series of gold catalysts supported on hydrotalcites also showed promising activity. Zhang’s group discovered that thiolated Au₂₅ nanoclusters supported on calcined ZnAl-HT showed complete 3-nitrostyrene conversion in a wide temperature window from 90 to 135 °C with 3-vinylaniline selectivity higher than 98%. The high selectivity was attributed to the ZnAl-HT support adsorbing nitro groups rather than
vinyl groups [282]. Follow-up research comparing MgAl-, ZnAl-, and NiAl-HT as catalyst support observed that the amount of basic sites were in the order of MgAl > NiAl > ZnAl, while the 3-vinylaniline yield followed the order of MgAl < NiAl < ZnAl, suggesting that medium to weak basicity of the HT surface might benefit the reactivity [283]. Because nitro groups were generally believed to readily adsorb on basic surface, the role of calcined HT is speculated to be adsorption sites for nitro groups, in synergy with the gold particles which activate hydrogen. Nonetheless, the choice of HT support used for nitrostyrene hydrogenation catalysts still request special attention, as another research indicated that the selectivity toward vinyl hydrogenation could be enhanced by changing HT composition [284].

8.2. Hydrogenation of Nitriles

Hydrogenation of nitrile is an important industrial route for amines production. To inhibit byproducts such as secondary or tertiary imines, bases such as alkali bases or ammonia are commonly added along with metal catalysts [285]. The inherent basicity of MgAl-MMO materials makes them potential candidates for the hydrogenation of nitriles to produce amines. Tichit and coworkers prepared a series of Ni-containing catalysts based on LDH precursors for catalytic hydrogenation of nitriles. They discovered that the introduction of Mg in LDH precursors was effective in decreasing the surface acidity of the catalyst, decreasing the adsorption strength of primary amines, and thus preventing further coupling reaction between amines and imines. [286,287]. Cao et al. formulated a core–shell catalyst by coating nickel-based nanocomposites with LDHs or layered double oxides (LDOs), to catalyze the selective hydrogenation of benzonitrile to N-benzylaniline or benzylmethylamine. It was found that the structural and acid-base properties of the LDH/LDO-coated nanocomposites could be switched by calcination or hydration according to the memory effect of LDHs [288]. Their following work on LDH-derived NiMgAl-MMO catalysts for the hydrogenation of benzonitrile also showed that the presence of strong metal–support interactions in NiMgAl catalysts could effectively inhibit the leaching or aggregation of Ni nanoparticles, accounting for the excellent stability of NiMgAl-MMO than the impregnated Ni/Mg$^{0.75}$Al$^{0.25}$ catalysts [289]. The strong metal–support interaction was also observed in another study, in which a highly stable and active Co$_2$P supported on HT was prepared and studied for the ammonia-free selective hydrogenation of various nitriles to corresponding primary amines [290]. Compared to Al$_2$O$_3$, SiO$_2$, or carbon support, HT support stabilized Co species and prevented their oxidation, leading to enhanced reactivity and stability.

9. Future and Prospect of LDH-Derived Catalysts for Hydrogenation and Hydrogenolysis

Hydrogenation and hydrogenolysis are important reactions particularly in the petrochemical industry, biomass upgrade and CO$_2$ conversion. In the past decade, researchers have carried out extensive research on the application of LDH-derived catalysts in hydrogenation and hydrogenolysis. As reported in numerous studies mentioned above, superior reactivity of LDHs and their derived catalysts undoubtedly demonstrates outstanding advantages over conventional catalysts in the following aspects:

(1) Various facile preparation methods, including co-precipitation, hydrothermal methods, ion exchange, urea hydrolysis, etc., could be used to prepare LDH-derived catalysts. These preparation methods are generally mature and well-established, facilitating the wide application and scale-up production of LDH materials.

(2) The appropriate amount and strength of acidity/basicity are crucial for the design of multifunctional catalysts, and these properties could be achieved with LDH-derived materials. LDHs and MMOs possess high concentrations of acid/base sites, which are key sites for adsorption and reaction. For example, as discussed in this review, acidic sites could catalyze deoxygenation reactions, while basic sites could interact with carbonyl groups or catalyze aldol condensation. More importantly, the acid–base properties of LDHs and their derivative materials can be finely modulated by controlling the main layer element composition, interlayer ion species of the LDH precursor, and other synthesis parameters.
(3) Metal particle size is crucial for the reactivity and selectivity of metal catalysts. In hydrogenation or hydrogenolysis reactions, metal catalysts with high metal dispersion will provide more active sites for dissociative activation of hydrogen and surface reaction to take place, resulting in increased reaction rates and decreased catalyst usages. For LDH-supported metal catalysts, hydroxy groups and interlayer galleries could promote the dispersion of metal particles. For MMO or IMC catalysts, because metal cations are uniformly distributed in the atomic level within LDH layers, the calcination of LDH precursors would generate MMOs with highly dispersed metal atoms. Therefore, LDH-derived materials become ideal catalysts for hydrogenation or hydrogenolysis due to their capability of generating small metal particles.

(4) For MMO or IMC catalysts, when LDHs are calcined at high temperatures, active metal atoms are immobilized in the metal oxide matrix. This interaction between active metals and oxide matrix results in the formation of strong metal–support interactions (SMSI). SMSI is important not only in effectively preventing particle aggregation or sintering of the active metal during reaction but also modifying the electronic properties and catalytic reactivity of active metals.

Although much progress was made in the field of hydrogenation or hydrogenolysis catalysis over LDH materials, several challenges remained as follows:

(1) The synthesis procedures of LDHs were studied thoroughly, but for LDH-derived catalysts more optimization and mechanism research on the preparation methods are still necessary. For example, for MMO catalysts, the relationship between preparation parameters (e.g., metal precursors, precipitation pH, crystallization time, calcination temperature, reduction temperature, etc.) and physiochemical properties is still vague or case-dependent.

(2) The structure of LDH-derived catalysts is also currently unclear, which deserves more characterization efforts or theoretical predictions. The surface composition of LDH-derived catalysts, the electronic and geometric interactions between metal nanoparticles and neighboring components, and the origin of acidity/basicity of LDH-derived materials, are largely unknown, leading to difficulties in studying structure–reactivity relationships and catalyst design.

(3) LDHs show a unique “memory effect” and good reversible topological conversion properties after heat treatment over a wide range of temperatures. Accordingly, how LDH-derived catalysts go through structure transformation during the reaction process also needs in-depth research. For biomass-related hydrogenation or hydrogenolysis research, this issue is important because water is often present either as reactant or product, which might lead to topological transformation of LDH derivates. More research on structure transformation of LDH-derived catalysts before, during, or after reaction will be valuable for a broader application of LDH-derived catalysts.

10. Conclusions

In this review, the applications of LDH-derived materials in the field of catalytic hydrogenation and hydrogenolysis are comprehensively summarized. LDH-derived catalysts in hydrogenation and hydrogenolysis could be categorized into three types: LDH-supported catalysts, mixed metal oxides, and intermetallic compounds, while different types of catalysts exhibit different structure and physiochemical properties. The tunability in composition and property of these LDH-derived materials makes them versatile in many applications. As demonstrated by numerous examples mentioned above, LDH-derived catalysts showed superior reactivity owing to their unique advantages. Therefore, based on the current research progress, it could be envisaged that research on hydrogenation or hydrogenolysis over LDH-derived catalysts will keep on increasing rapidly in the future, and industrial applications of LDH-derived catalysts will also be expanding due to their desirable characteristics.
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