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Chapter 1

Metal Nanoparticles as Emerging Green Catalysts

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Abstract

Green nanotechnology is defined as the technology applied for building clean technology by which one can reduce the potential risks of environment and also improve human health conditions. It is linked with the implementation of products of nanotechnology and its process of manufacturing. Green nanotechnology synthesizes new nanoproducts with improved properties in such a way that they can substitute some of the existing low-quality products. The main motive of developing new nanoproducts is to enhance sustainability and also to make them more environment friendly. In particular, nanoscale materials (e.g., nanoparticles) can be defined as those having characteristic length scale lying within the nanometric range, that is, in the range between one and several hundreds of nanometers. Within this length scale, the properties of matter are sufficiently different from individual atoms/molecules or from bulk materials. The primary objective of this chapter is to provide comprehensive overview about metal nanoparticles (MNPs) and its application as emerging green catalysts. This chapter contains six sections in total. Section 1 starts with a general introduction, recent progress, and brief summary of the application of MNPs as green catalyst. Section 2 reviews the preparation and characterization of supported metal nanoparticles for a wide range of catalytic applications. Section 3 presents the catalytic properties of supported metal nanoparticles. Section 4 describes briefly some of the most commonly reported supported MNPs in different green catalytic applications. Section 5 concentrates on our own results that related to the application of supported MNPs in catalysis. In this section, the oxidation of benzyl alcohol to benzaldehyde, the production of adipic acid from cyclohexane, the photodegradation of dyes using green route will be discussed. Finally, Section 6 describes the summary of main points and also presents an outlook of the application of MNPs in green chemistry.

Keywords: metal nanoparticles (MNPs), MNPs for environmental application, green catalyst, photocatalysis, MNPs characterization, oxidation reaction
1. Introduction to metal nanoparticles

Materials whose length lies within the nanometric range of one to several hundred nanometers are known as nanoscale materials or nanoparticles [1]. Matter at this scale is characterized by significantly different material properties than molecules, atoms, or bulk items [2]. Because of this, the field of Nanoscience was coined to address this relatively novel scientific field. Although the concept of this scientific area is not new, contemporary science is constantly discovering new and favorable applications for both the manufacture sector and academia, which has cast Nanoscience as a burgeoning and exciting field. The applications referred to are diverse, and range from environmental development to consumer products, and even therapeutic use (Figure 1).

![Figure 1. Potential application of nanoparticles in various fields.](image)

Nanoparticles can take many forms, including powder, crystal, and cluster formulae. Nanopowder is used to describe mixtures of fine powder, whereas ultrafine particle mixtures are described as nanocrystals [2]. Clusters can be further classified as nanoclusters if they have a narrow size distribution in the range of 1–10 nm and a minimum of one dimension. Prior to the 1700s, nanoparticle use was limited to high-quality glass, pottery, and tile production. In the mid-nineteenth century, Michael Faraday began to explore nanoparticle stability. At the beginning of the twentieth century, Mie conducted research into nanoparticle color properties.
Shortly thereafter, Richard Zsigmondy received the 1925 Nobel Prize in Chemistry for his work on colloidal suspension of gold nanoparticles using ultramicroscopic techniques.

Metal nanoparticles (MNPs) are deemed to be likely candidates for catalysis due to their relatively large surface area per volume or weight unit as compared with bulk metal, meaning heterogeneous MNPs catalysts typically function on metal surfaces [3]. A variety of techniques are used to develop supports for industrial metal catalysts, which are frequently inorganic and characterized by highly dispersed metals and voluminous surface area. Metal particles tend to be ensemble affairs, consisting of more than one particle in a range of forms and dimensions. Polyhedral particles are formed from the interaction of MNPs with the inorganic supports. Silica gel, activated charcoal, and alumina, among others, are examples of transition metals that are formed from metal particles distributed onto inorganic supports. These are superior to metal powders for numerous reasons, including their stable thermic capacity, diminished cost, greater surface area, and efficacious use of MNPs in the form of wide dispersion.

The electrochemical makeup of particularly dimensioned MNPs allows them to be used as catalysts. As compared with bulk metallic catalysts, greater surface area and measured edge step ratio result in higher outputs. MNPs also are well suited to stereoselective synthesis. MNPs have been increasingly examined in recent years due to their catalytic properties in a range of contexts, for example, as a surfactant for water-soluble polymer, resins, vesicles, and so on. Other examples are the reactions including Fischer-Tropsch, isomerization, and hydroformylation, etc., which also use nanoparticles such as Rh, Pt, Ir, Au, and Pd as catalysts [4]. The size of the metal particles holds great influence over the catalytic abilities of the metal. Additionally, if one looks to the use of gold nanoparticles in catalysis, the size of the gold particle is directly linked to catalytic activity and/or selectivity. Gold particles with diameters less than 10 nm are deemed to be very active catalysts, while such activity considerably diminishes as the Au diameter increases, and also virtually disappears when the diameter is measured in micrometers range [5, 6].

MNPs are functionalized in many situations as they typically have greater stability in solutions as compared to nonfunctionalized metallic nanoparticles. This is because the nonfunctionalized particles present with a reduced surface area and thus lower catalytic ability due to their tendency to aggregate while in solution. Although ligands can be used to avoid aggregation, they have the consequence of altering catalyst functionality. Nanocatalysts can also be functionalized using polymers and oligomers. Typically, catalytic nanoparticles have been implemented in tandem with polymer matrices; otherwise, they were detained on solid supports [7]. The petrochemical industry has a long history of utilizing supported metal catalysts, particularly Pt, which is used for naphtha reforming. However, new techniques that employ electrostatic interaction forces are now being created that will enable stable catalytic aggregate of a preferred form [8]. Multilayer polyelectrolyte films is formed through the adsorption of metal ions (Ag+, Pt2+) [9].

MNPs are primarily defined by their size; so, it follows that size is the real driving force of their catalytic capabilities. Because metal particle surface area is inversely proportional to squared nanoparticle diameter, as particles decrease in size, the surface area increases. This decreasing size results in greater catalytic activity, as all chemical reactions occur on the
catalyst surface [10, 11]. The rate of reaction is also influenced by particle size in case of some chemical reactions. In these instances, slower reactions are caused by reduced nanocatalyst size. For example, when using Pt nanocatalysts for the generation of photochemical hydrogen, it is ideal to use catalysts with 3 nm particle diameters. Any other size, greater or less, will slow down the chemical reaction [12]. Catalyst selectivity is also affected by size. For example, when creating monoene in cyclopentadiene partial hydrogenation, a size of <2 nm is considered ideal. However, the particles having larger size will alter the reaction's selectivity due to the Pt nanoparticles' steric effects [13].

In addition, material arrangement of nanocatalysts also plays a key role in catalytic activity, wherein the addition of other metals can result in added value. The ligand and ensemble effects are most often cited as the cause of this effect. Because of the redox potential of metals, the reaction is also impacted by nanocatalysts' core-shell construction [14]. As an example, the ratio of Pd/Pt = 4 is considered to be the ideal nanoparticle ratio for high performance during the process of selective partial hydrogenation of 1,3-cyclooctadiene to cyclooctene. However, these bimetallic nanoparticles will experience diminished catalytic activity if such ratio is altered. The ligands effect resulting from the incidence of Pt at the particle cores means the bimetallic catalysts have greater activity levels as compared with monometallic Pd catalysts [15].

The catalytic performances of supported MNPs are dependent on catalyst stability, which can be proven when considering the relationship between MNP size and catalytic activity. However, agglomeration means naked colloidal nanoparticles, which are already thermodynamically unstable, are further unstable in the long run. In order to prevent such agglomeration, these particles must be stabilized for the long term, which can be achieved in a number of ways. Two methods would be supporting them on a solid surface or coordination with ligands/anionic species [16]. Metallic nanocatalysts can be stabilized in either aqueous or nonaqueous solutions using polymers, as is the case with the use of polystyrene-b-poly-4-vinylpyridine being used to ensure Pd is stable during a Heck reaction in which a styrene and 4-bromoacetophenone coupling takes place [17].

2. Synthesis and characterization of metal nanoparticles

2.1. Synthesis methods [1, 2]

Metal particle synthesis typically takes place in one of two ways: top-down or bottom-up, as shown in Figure 2. The top-down approach uses an external force to pressure bulk materials, eventually causing these materials to break down into smaller components by means of mechanical, chemical, or some other energy sources. A bottom-up approach takes place in a reverse tactic, growing precursor particle size by using chemical reactions to combine atomic or molecular species. It should be noted that the top-down approach is considered to be a physical method while the bottom-up approach is chemical, although both approaches can be applied in a range of states, including liquid, solid, gas, supercritical fluids, or vacuum. The final desired outcome and manufacturer must play into the choice between the two methods.
Because size and shape of the metal particles is the most vital aspect to be altered for improved application performance, it is a vital aspect to take into account for catalysis as an example. This section will address some selected methods for fabricating metal nanoparticles, including physical and chemical methods.

2.1.1. Chemical methods

When it comes to synthesizing metal nanoparticles from a range of source materials, chemical methods have become a reliable technique. They are relatively simple and are implemented under mild conditions. The following section will examine the production process for isolated nanocrystals dispersible in solvents (sols).

2.1.1.1. Chemical reduction

The most ancient and widely used chemical method for synthesis of nanoparticles is the reduction of metallic ions in solution. In ion reduction method, metal ions are reduced by providing some extra energy and using the different types of chemical reductants. The provided energy is used to decompose the material, and usually, photo energy, electricity, or thermal energy is used. It is the most frequent chemical method used for the production of stable metallic nanoparticles. Turkevich used this method to produce spherical nanoparticles of gold by reducing the gold hydrochlorate solution with sodium tricitrate [18]. Nanoparticles of gold can also be produced by reducing the solution of chloroauric acid; this reaction reduces $\text{Au}^{3+}$ ions to $\text{Au}^{+}$ and further into $\text{Au}^0$. Brust method and Perrault method were also used to
produce metallic nanoparticles; the main theme of both these methods is the same as in Turkevich method with different reducing and anticoagulant agents.

2.1.1.2. Coreduction of mixed ions

Coreduction of mixed ions is similar to chemical reduction in many ways. However, this method is mainly used for preparing bimetallic nanoparticles. The preparation of the colloidal dispersions of bimetallic nanoparticles containing gold can be prepared using chemical methods [2]. Coreduction is the simplest preparative method of bimetallic nanoparticles. In this method, metal ions of two or more metals are usually reduced by suitable reductants (e.g., citrate) [18]. The average diameter of bimetallic nanoparticles depends on the metal composition.

2.1.2. Physical methods

Physical methods require solid material to be evaporated into supersaturated vapor, which supports the homogenous nucleation of nanoparticles in order to prepare MNPs [1]. This also allows particle size to be temporarily controlled either by introducing gas molecules that collide with the particles, or by momentarily halting the evaporation source. Because growth occurs over a matter of milliseconds or seconds, experimental parameters must be securely controlled. Recent years have seen the development of several particular methods that can be categorized according to their use of solid or liquid (vapor) precursors, as well as the energy source employed.

2.1.3. Other methods

In this section, some additional methods for preparing nanoparticles are reviewed. These methods include both chemical and physical methods.

2.1.3.1. Vapor synthesis

In order to create conditions that support particle nucleation, a hot wall reactor is used on material, and other variable factors (pressure and temperature) are adjusted during the vapor phase. The vapor synthesis model is favored for multicomponent or bimetallic nanoparticles. This process has been recently used by Schmechel to translate organometallic yttrium and europium to europium-doped yttria nanoparticles [19].

2.1.3.2. Sputtering

Inert gas in the form of high-velocity ion is applied in this technique to vaporize material from a solid surface. The application results in the solid surface ejecting atoms, and is typically done in a vacuum state. It is also possible to deploy electrons in the place of atoms [20]. This technique is preferred for its ability to create nanoparticles with a composition matching that of the targeted material [21].
2.1.3.3. Laser reactors

This technique uses laser beams to heat a substrate in the presence of an inert gas. Laser beam energy is absorbed by the substrate during the vapor phase, resulting in direct warming. As a result, temperature increases due to collision, and despite the fact that inert or carrier are not directly heated. The dropping temperature of inert gas causes supersaturation and prompts the formation of nanoparticles on the reactor wall [22]. Laser pyrolysis is typically carried out with ruby laser and CO\textsubscript{2}-type lasers. The fact that there is no need to use a heat wall is a major benefit of this method. This avoids contamination, which can otherwise be introduced by the heat wall, making this method more efficient than sputtering methods. However, the technique has drawbacks as well—the use of laser increases costs [23].

2.1.3.4. Flame reactors

Nanoparticles can also be produced using flame, whose heat initiates a reaction resulting in condensable monomers that are frequently in agglomerate form. The flame reactor method is considered cost-effective [24]. For instance, TiCl\textsubscript{4} can be heated and oxidized in the presence of flame, resulting in the nanoparticles of TiO\textsubscript{2}. This is just one example on the use of this method. Small modifications allow flame reactors to be used for complex and free agglomerate outputs as well [25].

2.1.3.5. Plasma reactors

Nanoparticle production can be initiated using plasma as an energy source. This is because plasma temperature is high enough to isolate atoms and radicals from the substrate, which results in nanoparticles establishing themselves on the cooling gases. Dc arc plasma, dc plasma jet, and radio-frequency (rf) induction plasma are the most frequently used plasmas in this method, which can be used to generate multicomponent nanoparticles [26, 27]. It is possible to speed up production by introducing cold gas to plasma-formed vapors, but this can cause destruction of nanoparticle homogeneity [27].

2.1.3.6. Wire electrical explosion

Heat and vapors are produced in this method by passing a high current through a metallic wire. It is useful when dealing with metals that are difficult to vaporize in a furnace due to their high boiling point (Si, C) [28]. The method has previously been used to create 2–4 nm Si nanoparticles using Ar inert gas as a carrier. Reaction and carrier duties are carried out by reactive gases to form binary metallic and multicomponent nanoparticles [29].

2.1.3.7. Expansion–cooling

Abrupt expansion and cooling are used to stimulate homogenous nucleation by passing condensed gas through a nozzle, resulting in gas expansion and temperature reduction. Particle size is then able to be altered using pressure change: bigger sizes are achieved using higher pressure. For example, N\textsubscript{2} gas in a subsonic nozzle will result in 100 nm particles if 2
bar pressure is applied. However, using 0.75 bar pressure under the same conditions produces 5 nm particles [20].

2.1.3.8. Electrospray systems

This is the simplest means of creating nanoparticles and requires a small droplet of dilute solution to be evaporated. The intended matter is made into diluted solution and dried in small droplets of 1 μm, producing nanoparticles. However, it is difficult to create this very small size of droplet required, and hence there is an added challenge that the solvent can introduce some impurities [30].

2.2. Characterization methods

Nanoscale materials have a unique set of properties, making them suitable for a variety of applications and resulting in them being a focal point for a range of researches. Deep understanding of the properties of metal particles at nanoscale level is necessary to understand the nature of active sites, which in turn can help to find and tune the key performance signs. Several techniques exist that are designed to characterize nanoparticles, although none of these are able to supply full information regarding investigated materials. For this reason, several techniques must be used to categorize each sample in order to understand the size, structure, and catalytic properties. Figure 3 depicts various methods of analyses for MNPs.

<table>
<thead>
<tr>
<th>Characterization Method</th>
<th>Information obtained</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dynamic light scattering (DLS)</td>
<td>Size and size distribution of MNPs in solution</td>
</tr>
<tr>
<td>UV-Vis spectrum</td>
<td>Formation of colloidal MNPs (Plasmon band)</td>
</tr>
<tr>
<td>X-ray diffraction (XRD)</td>
<td>Crystal structure and size, chemical composition</td>
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<td>Transmission electron microscopy (TEM) Scanning electron microscopy (SEM)</td>
<td>Size and Morphology of MNPs</td>
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<tr>
<td>Energy Dispersive X-Ray (EDX)</td>
<td>Element and distribution of MNPs</td>
</tr>
<tr>
<td>Scanning tunneling microscope</td>
<td>Size and structure of MNPs</td>
</tr>
</tbody>
</table>

Figure 3. Summary of some selected characterization techniques of MNPs.

Common means of characterizing MNPs include transmission electron microscopy (TEM) and scanning electron microscopy (SEM). Other common techniques include X-ray photoelectron spectroscopy (XPS), UV spectroscopy, atomic force microscopy, powder X-ray diffractometry (XRD), and dynamic light scattering (DLS). UV spectroscopy and Fourier transformed infrared spectroscopy (FTIR) methods would be combined to examine gold and palladium nanoparticles.
cles, as described elsewhere [31, 32]. Conversely, silver nanoparticles would be examined using TEM, high-resolution TEM, and selected area diffraction pattern (SAED) [33]. Finally, X-ray diffraction, SEM, and FTIR would be combined to examine magnetite, Ag, zinc, and Au nanoparticles [33–35].

3. Catalytic properties of supported metal nanoparticles

Supported metal nanoparticles (SMNPs) are used today in several application fields such as sensors, nanoelectronics, medical-technical applications, and in heterogeneous catalysis [36, 37]. In general, SMNPs consist of metal nanoparticles (MNPs) from transition metal or noble metal pools that are deposited on high surface area supports (150–1200 m$^2$/g). In general, the metal load of supported metal catalysts is generally in the range of 0.5 up to 10 wt. % (even higher in rare cases). MNPs are often in a size range of roughly 1–30 nm. Such particles reveal chemical and physical properties that are intermediate between those of a single metal atom and a bulky metal particle of a size much larger than 100 nm. However, careful handling is important, that is, once activated, most of these SMNP solids are pyrophoric by contact with oxygen.

The activity of supported metal catalysts in heterogeneously catalyzed reactions is mainly determined by the shape and size of the metal species (i.e., MNPs) deposited onto the support, because only surface metal atoms might take part in a chemical reaction [37–39]. Heterogeneous catalyst performance is very sensitive to particle size, because the structure of the solid surface and the electronic properties may significantly change depending on the size of the MNPs [37, 38]. For example, the activation energy for CO dissociation and the heat of adsorption for CO alter with changing size of MNPs. These changing properties affect its performance in several CO-converting reactions like Fischer-Tropsch (FT) synthesis for the manufacture of hydrocarbons from gas synthesis. Supported Au-containing catalysts show similar size effects with respect to applied supports [40, 41]. Au nanoparticles on alumina smaller than 4 nm show complete conversion in ethylene glycol oxidation, but increasing nanoparticle diameter above 5 nm leads to significant drop to roughly 40% conversion only [40].

Therefore, one crucial property of supported heterogeneous MNP catalysts is that they contain the metals as finely dispersed particles on the support surface, in general [38, 42, 43]. Incidentally, the term dispersion describes the degree of metal distribution (Figure 4); a dispersion degree of close to 1 means that all offered metal atoms exist in an atomic dispersed monolayer at the surface, whereas a dispersion degree of almost 0 says that only one large particle is present [44, 45]. Thus, the dispersion degree is inversely proportional to the MNP size. The dispersion degree might be determined by chemical methods such as hydrogen or CO chemisorption; a nice overview on CO chemisorption on metal particles is given by Hammer et al. [46]. However, one has to know adsorption chemistry well to draw the right conclusions from experimental data; for example, CO can be adsorbed on Pt surfaces linearly, bridged or multibound, as proven by different bands obtained from FTIR spectra [47]. Thus, the number of CO atoms adsorbed on a nanoparticle surface does not necessarily mirror the number of
surface atoms and the dispersion degree. Nowadays, DFT calculations and several adsorption models help to refine experimental data [48]. Small-angle X-ray scattering (SAXS), X-ray diffraction, and TEM can also be applied; in addition, several in situ studies can be included for characterizations [37, 49]. The latter needs some more intense sample preparation and examination efforts to get statistically reliable conclusions.

**Figure 4.** Well-dispersed metal nanoparticles (MNP) versus a large metal particle on a support surface.

Besides MNPs’ size and shape, the catalytic properties of such particles might be influenced by a second metal, the so-called bimetallic catalysts. The addition of a second metal often leads to alloys showing quite different electronic and also structural properties. Some examples are the formation of PtNi or Pt-bimetal SMNPs for electrocatalytic applications [50, 51] or AuNi SMNPs for steam reforming of hydrocarbons [52]. Moreover, support properties strongly influence the catalytic performance of SMNPs. Matsumura and Nakamori describe in detail the effect of various supports on catalytic properties of such nickel catalysts in the steam reforming of methane (SRM) reaction [53]. Au nanoparticles (<4 nm) on alumina reveal full conversion in glycol conversion, whereas Au nanoparticles on carbon need a size of 7–8 nm to reach complete conversion [40]. Support-metal interaction is studied in detail for several catalyst systems and also reviewed in Ref. [54].

Mainly hydrogenation (dehydrogenation), oxidation, automotive off-gas cleaning, and C-C coupling reactions run on such SMNP catalysts [42, 43, 55]. Most of these reactions proceed in the gas phase, and various examples of liquid phase or trickle bed reactions are also known. In particular, in the latter cases, loss of active MNPs by leaching might be a catalyst deactivation reason. Further deactivation reasons are due to MNPs sintering by high reaction temperatures or easy movability on a support surface. Furthermore, MNPs might deactivate by accumulation of products or oligomers and coke-like products on their surface. In general, the latter can be easily regenerated by hydrogen or air (oxygen) treatment in case of oxidic carriers with subsequent reduction. Au, Pd, Rh, Ru, Ni, or Ag are the mostly widely used MNPs for the above-mentioned transformations [55]. In general, various oxidic supports, such as diverse aluminas, silicas, titanias, etc., and several carbon materials, are usually used as supports for the MNPs. White et al. recently summarized various synthesis protocols in a concise review [55]. Impregnation, coprecipitation, and deposition-precipitation are some of the classical chemical synthesis routes. Precursor compounds have to be first calcined under air and second reduced by hydrogen or hydrides to receive MNPs. Physical synthesis routes include the application of microwaves, sonic waves, or supercritical fluids. Moreover, there are several protocols to deposit pre-prepared colloidal MNPs onto the surface of various supports [56]. This is a very interesting approach, because colloidal particles can be synthesized very uniformly with respect to size and shape [37, 56].
Supported metal catalysts are applied in large scale in several industries such as refining of petroleum, hydrogenation of carbon monoxide, hydrogenation of fats, and many other processes [43, 55, 56]. They are also the heart of automotive catalysts to clean automobile exhaust gases [57]. Such catalysts and the significance of a high dispersion are known for over hundred years, in particular, from various hydrogenation reactions. The hydrogenation of carbon monoxide to methane over supported nickel catalysts was discovered in the beginning of the last century by Sabatier and Senderens [58, 59]. This reaction is known today as “Sabatier reaction” or methanization, in general. It is applied in large scale in industry to remove traces of CO from ammonia synthesis gas, but it also gains importance in energy-storage efforts by converting solar energy via hydrogen to storable methane [60]. In this process chain, CO$_2$ reacts nearly quantitatively with hydrogen to methane over Ni or NiRu containing catalysts at about 300–350°C, with H$_2$:CO$_2$ feed ratio of 4 and slightly increased pressure up to 10 bar [60, 61].

Paul Sabatier received together with Victor Grignard the Noble Prize in Chemistry in 1912 for his method of hydrogenating organic compounds in the presence of finely disintegrated metals, whereby the progress of organic chemistry has been greatly advanced in recent years [62]. The explanatory statement explicitly points to the necessity of the “finely disintegrated metals” that we know today as metal nanoparticles.

One of the most common hydrogenation catalyst formulation is nickel on oxidic (e.g., alumina, zirconia, ceria, titania, or silica) [55, 63, 64] or carbon [65, 66] supports. As mentioned above, such catalysts are normally used as methanization catalysts, and other applications are also in common use such as catalysts, for example, for hardening of oils and fats in food industry, dry reforming of methane to manufacture syngas by using CO$_2$ [67, 68], or in petrochemical industries and refinery. In the latter case, they are of interest in the hydrogenation of various aromatic compounds to reduce their content in fuels. Nickel-containing catalysts are also in large-scale operation with respect to SRM reaction to generate hydrogen and/or syngas (CO, H$_2$) [53, 69]. The SRM reaction is carried out in general over Ni/alumina catalysts at around 800°C. Additional hydrogen can be manufactured by the water gas shift (WGS) reaction converting CO and water to CO$_2$ and H$_2$. The WGS reaction is industrially carried out using supported high-temperature shift FeCr (350–500°C) and low-temperature shift CuZn catalysts (180–250°C), respectively [70, 71]. Nowadays, this reaction gains much importance for ion fuel cell applications too [72]; mainly nonnoble metals such as Cu or Ni on ceria are under development [71].

Nickel-containing catalysts play an increasing role in chemical transformation of biomass-derived feedstock. Song et al. studied Ni on various carbons or MgO supports for depolymerization of lignosulfonate and identified three Ni functions being (i) the active sites for hydrogenolysis of C–O–C bonds; (ii) as active sites for hydrogenolysis of C–OH bonds at side chains to alkane chains, and (iii) as convertors for sulfonate to H$_2$S [73]. In a detailed study on hydrodeoxygenation of phenol as model compound and pyrolysis oil as a real feed, we recently could show the potential of bimetallic NiCo NPs supported on zeolite supports [74].

Further examples of the industrial application of SMNP catalysts are FT synthesis of hydrocarbons over Co, Fe, Ni, Ru catalysts [75, 76] or the synthesis of ethylene oxide from ethylene and oxygen over Ag/alumina [77, 78]. Another good example is the use of sulfided CoMo and
NiMo catalysts in refinery industry for the removal of sulfur from different distillation cuts by special hydrodesulfurization (HDS) units [79]. Mainly alumina is used as support; however, improvements in deeper HDS are reached by a variation of supports (silicas, zeolites, amorphous aluminosilicates, etc.) or the addition of further (noble) metals to the conventional catalyst composition [80, 81].

Besides these applications in well-studied industrial fields, novel SMNP materials are manufactured in recent years that might lead to new catalyst developments with respect to synthesis procedures, catalyst performance, and possible applications. Novel support materials like carbon nanotubes [82], graphene oxide [83], or carbon nitrides [84] might lead to novel interactions with MNPs resulting in novel catalysts. Jin et al. recently reported on the use of mesoporous metal oxide microspheres as support for Pd NPs and their use in reduction of nitro compounds [85]. Increasingly, noble metals are replaced by transition metals, or their alloys lead to cheaper catalysts, improved performance, NP stability, and/or new applications [37, 60, 86].

4. Green catalytic application of metal nanoparticles

Even though most of the catalysts are made by mixing and shaking some components, their surface structures are not yet well controlled. For any catalyst, it is desired to achieve long life cycle, 100% selectivity, and low energy consumptions, and these properties can only be gained by controlling their size, shape, electronic structure, and chemical and thermal stabilities of catalyst particles [87]. In the case of dealing with catalysts in the nanoscale level, selectivity ratio can be adjusted by precisely controlling the size of nanoparticles. Hence, precious metallic catalysts are usually used in the form of nanoparticles to improve their selectivity.

Nanocatalysts are used in chemical processing industries due to their lower energy consumption, environment-friendly behavior, with an improved economy. For instance, nanocatalysts of NiO with novel aluminum oxide support are used in the production of syngas and bio-oil from pyrolysis of biomass. Nanocatalysts improved the quality of syngas by decreasing the fraction of carbon monoxide in the syngas; it also reduced the proportion of tar in the product [88, 89].

Biomass energy is fulfilling about 15% of total energy requirement of the world; the broad range of organic materials included wood, crops, organic wastes (animal dung, municipal wastes, etc.), which are used for the production of biomass energy. And mostly, these materials are produced from the process of photosynthesis; according to the study of Luo and Demirbas, approximately 720 tons of biomass are produced per year [90, 91]. There are three main types of reactions for the conversion of biomass into fuel energy, such as combustion, pyrolysis, and gasification. Pyrolysis technique is preferred on other techniques due to its product named as pyrolytic oil or bio-oil. High moisture content, long-chain hydrocarbon, and low hydrogen-carbon ratio make it difficult to burn. Therefore, mainly biofuel is not directly used as fuel. Aravind [92] and Malik [93] studied the effect of nanocatalysts on the processing of bio-oil into syngas and concluded that nanoparticles with high surface area are the best alternatives of
conventional catalysts to improve the yield of syngas. Tar is an undesirable by-product in syngas production due to its high boiling point, and hence it blocks the filter and pipes. Tar is also a poison to the catalyst [94]. There are two methods to control the production of tar: one is treatment within gasifier and the second one is hot gases treatment after the gasifier. The second method is most economical, and hence it is only used in industries. Nanocatalysts of sodium, potassium, and calcium are used for hot gases treatment of effluent to reduce the amount of tar [95]. Catalytic reforming of tar by using metallic nanocatalysts is preferred because there is no need for any additional energy, and that is the reason why nanocatalysts are called green catalysts. This catalytic reaction also includes steam reforming, hydrocracking, and hydroreforming [96, 97].

In the Fischer-Tropsch synthesis of green diesel, nanocatalysts of iron and Co of size 10–15 nm are used in slurry reactors to improve the production of high molecular waxes. These waxes then hydrocracked to generate green diesel [98, 99]. Fischer-Tropsch’s is an important method to convert nanopetroleum feedstock such as coal and biomass into clean diesel, which can be used as a fuel. Uses of nanocatalysts dramatically decrease the cost of this process as reported elsewhere [100].

FT syntheses technology produced high-quality ultraclean fuel with low aromatic content and zero sulfur. The product mostly consists of the mixture of linear and branched hydrocarbons [101]. Two types of techniques are adopted for FT synthesis, but the slurry bubble column techniques gain more interest due to its low pressure and stable temperature [102]. Generally, the Fischer-Tropsch synthesis follows the ASF distribution mechanism, but this distribution is unselective in this case. Later on, the novel Fischer-Tropsch catalysts are developed; the novel FT nanocatalyst containing Co with porous silica support is more selective toward hydrocarbons of C_{10}–C_{20} (diesel) [103]. Study of Kang et al. [99] showed that the nanocatalyst of Ru is more favorable in the high moisture content, but the only issue they have is low activity. This issue was solved by using carbon nanotube supported ruthenium catalyst, which gives excellent selectivity for diesel hydrocarbons and also high activity of the catalyst [99, 104].

Solid catalysts of Aluminum dodecatungstophosphate (Al_{0.9}H_{0.3}PW_{12}O_{40}) or nanotubes of Zn_{1.2}H_{0.6}PW_{12}O_{40} are used for esterification of waste cooking oil to produce biodiesel; these catalysts improved yield from 42.6 to 96%. It is due to improved surface area of the nanocatalysts from the bulk conventional catalyst [99]. Animal and used cooking oil are alternative sources of diesel fuel to run the vehicles [105]. However, high water content and the high ratio of free fatty acid make them unfavorable for the production of diesel fuel. Sulfuric acid with a large quantity of methanol at high temperature and pressure reduces the quantity of free fatty acids. High pressure and high temperature increase the diesel fuel production cost [106]. According to the study of Li et al., ZnPW nanotubes can be used to reduce free fatty acid at lower temperature and pressure [107].

MNPs are also used in fossil fuel sector to improve the reaction; selective hydrogenation, paraffin hydrogenation, naphtha reforming, and hydrodesulfurization are some of the major units where nanocatalysts are used in oil refineries. Nanoparticles hexanethiol monolayer-protected palladium of size 1.5 nm are used to improve energy consumption of catalytic
combustion of JP-10 activation fuel. These nanoparticles decrease the ignition temperature to 240°C [108, 109].

Meurig et al. [110] studied the bimetallic nanocatalysts of size range 3–30 nm with nanoporous silica support in single-step hydrogenation and found that this catalyst greatly facilitates the separation of product from reactant, with easy recycling at low temperature and pressure conditions.

Hydrogen gas can be used in fuel cell to generate green energy; now, almost 95% of hydrogen is produced by using partial oxidation and steam reforming of hydrocarbons. Both oxidation and steam reforming produce carbon dioxide, which is a major constituent of global warming [111]. Catalytic reforming of green biofuel such as ethanol can be a green way for the production of hydrogen gas; ethanol is usually available with high water content; the water present in ethanol and other impurities deactivate the conventional catalysts. Nicolas et al. [112] worked on the designing of nanocatalysts for the production of hydrogen gas by using ethanol and concluded that nanocatalyst rhodium metal with alumina support gives the highest yield of hydrogen gas with a better protection from water. Rhodium catalyst produces cleavage between carbon and hydrogen, and this is the main reason of C–C bond rupture. Hydrogen in fuel cell gives 50–60% efficiency, while in diesel engine only 20–25% fuel is converted into useful energy [113, 114].

In brief, MNPs are used to increase the selectivity and activity of catalyst, which resulted in less energy consumption and high yield. In general, the majority of the chemical reactions obtain on the surface of the catalyst and hence with decreasing size of particles there is an increase in catalytic activity. Slower reactions are caused by continuously decreasing the size of the metal nanoparticle catalysts whereas increasing the size of the catalyst will decrease the rate of reaction. Nanocatalysts are energy-efficient; they reduce global warming by reducing the required energy for any catalytic process with less poisoning ratio. They also decrease the chemical wastes; all these factors resulted into an improved economy and safer environment.

5. Own results

The nature of the support has great influence over MNPs’ catalytic performance. This is because the support allows for efficient pathways to the target outcome, which compounds the impact of particle size stabilization when it comes to small MNPs’ growth. Supports must have a strong metal-support interaction (SMSI), a high surface area, and incidence of active sites that can contribute to the reaction process [115]. This section investigates the catalytic application of MNPs, particularly for the oxidation of cyclohexane to adipic acid, the oxidation of benzyl alcohol to benzaldehyde, and the photodegradation of dyes. These reactions can be evaluated through the technique of placing MNPs on various kinds of metal oxide supports. The supports are used to afford high metal surface areas as well as stabilize the small MNPs characterized by high dispersion degrees [116]. Catalytic performance of heterogeneous reactions can also be affected by the support material characteristics, type of catalyst preparation method, degree of metal loading, and particle size. Specifically, supported noble metals
(e.g., gold) will have catalytic properties highly dependent on the type of support material at hand [116]. One example notes that acidic supports typically augment electron deficiencies on noble metals, when compared to basic supports. Strong metal-support interaction (SMSI) is demonstrated by metals supported on reducible oxides (e.g., TiO$_2$ and CeO$_2$) [117]. Robust interaction between the metallic components results in considerably higher catalytic activity compared to the catalysts having only one metal. This is in opposition to relatively inert irreducible oxides (e.g., Al$_2$O$_3$, SiO$_2$). This section will examine five kinds of supports, taking into consideration the benefits derived from metal oxide supports. These different kinds have different impacts in terms of particle size, dispersion, and performance. The oxidation of benzyl alcohol and cyclohexane were both tested with these five supports described below, and the reactions were performed in the liquid phase. In addition, evaluation of gold nanoparticles supported on TiO$_2$ and ZnO will also be evaluated photocatalytically. The outcomes of these examinations are detailed below.

5.1. Oxidation of benzyl alcohol to benzaldehyde

Green chemistry emphasizes the importance of using oxidation to formulate chemical intermediates and fine chemicals with high selectivity [118], as is the case with forming benzaldehyde (BAI) from benzyl alcohol (BA) oxidation (Scheme 1). The resulting solution is frequently used in pharmaceutical, agrochemical, and perfume industries. Reaction of benzyl alcohol with excessive ammonium permanganate or potassium is possible in aqueous acidic medium to form benzaldehyde (BAI). However, this method has significant toxic by-products formation, and hence it is not really environment-friendly. Efforts have been made to amend this process by oxidizing benzyl alcohol using a green oxidant in the presence of organic solvent, such as H$_2$O$_2$ or O$_2$, and applying various catalysts such as Pd/C, Pd(II) hydrotalcite, Pd-Ag/pumice, Ru-Co-Al hydrotalcite, Ni-containing hydrotalcite [119], but these techniques are still not considered to be environmentally acceptable due to the solvent use. Solvent-free oxidation is possible, using tert-Butyl hydroperoxide (TBHP) over MnO$_2$-exchanged hydrotalcite and a transition metal containing layered double hydroxides and/or mixed hydroxides. However, this is not a clean process as the TBHP produces tert-butanol. To be considered clean and environmentally friendly, there must be no solvent used, and the oxidant ought to be molecular oxygen (i.e., clean and inexpensive). Prati et al. successfully demonstrated such a process using Au/C catalysts to selectively oxidize alcohols and polyols [120]. According to Rossi et al., gas-phase oxidation of volatile alcohols into aldehydes and ketones is possible using Au catalysts [121]. A similar process oxidizing glycerol to glyceric acid using Au/graphite catalysts was carried out by Carretin et al. [122], and was deemed to be fully selective of the target output. This high selectivity of Au/CeO$_2$ was further proven by Abad et al. [123]. In case of benzyl alcohol oxidation to benzaldehyde, Enache et al. used supported Au on TiO$_2$ catalyst [124]. The same researchers also investigated the selective oxidation using Au-Pd alloy particles supported on TiO$_2$ [125]. In the end, the most efficient and environment-friendly means of oxidizing benzyl alcohol to benzaldehyde uses molecular oxygen. The technique also uses separable and recyclable supported gold nanoparticles over various metal oxide supports to provide high selectivity of the desired product, benzaldehyde.
For the past several years, the oxidation of benzyl alcohol is being investigated specifically looking at the impact of different supported MNPs (such as Au nanoparticles) on various metal oxide carriers [120, 125]. In the present study, we have used different oxide supports (CaO, MgO, ZrO₂, Al₂O₃, TiO₂) for AuNPs that were prepared with impregnation techniques [126]. A range of spectroscopic and microscopic techniques was deployed to characterize these catalysts and gather data regarding their individual properties. Catalytic performance of the AuNPs was tested on solvent-free oxidation of benzyl alcohol with oxygen (5 bar) as an oxidant in a Parr autoclave reactor at a reaction temperature of 140°C. **Figure 5** demonstrates that Au catalyst performance is strongly dependent on the kind of applied support. Although BAI is the primary output, some by-products were also produced, including small amounts of benzyl benzoate, benzoic acid, and acetal. Benzyl benzoate occurs because of the additional esterification reactions due to benzoic acid and benzyl alcohol. Benzoic acid is a natural occurrence when benzaldehyde over-oxidizes. Finally, acetal is produced when benzyl alcohol does not react and instead forms hemiacetal, which eventually forms acetal through protonation and deprotonation reactions. Acetal occurred in this instance only to a smaller extent, which is however negligible in this case. The most active catalyst found is TiO₂-supported AuNPs (at

**Scheme 1.** Oxidation of benzyl alcohol with O₂ using Au catalyst.

**Figure 5.** Influence of support on oxidation of benzyl alcohol to benzaldehyde over 1% Au/M catalysts (M = MgO, CaO, ZrO₂, TiO₂, Al₂O₃). Reaction conditions: 30 ml BA, 0.15 g catalyst, 140°C, 5 bar O₂, 4 h (X, conversion; Y, yield; S, selectivity).
X-BA = 81%), while the least active is MgO supported (at X-BA = 16% and S-BAI = 17%) (Figure 5). The enhanced performance of the former is due to several factors, including high dispersion of Au, smaller Au size, and comparatively greater surface enrichment of Au. These are completely opposite in the case of poor performing catalyst. TEM and XPS have been used to validate these claims.

5.2. Oxidation of cyclohexane to adipic acid

There are important industrial implications for selective oxidation of dicarboxylic acids from cycloaliphatic hydrocarbons, such as forming adipic acid (AA) from cyclohexane (CH) (Scheme 2). AA is a vital component when producing polyamides (such as nylon), polyesters, plasticizers (such as PVS), polyurethanes, and carpets. It is also used in the pharmaceutical and insecticides manufacturing processes [127], and has extensive applications for food products and medicines.

Commercial production of AA involves a two-stage process. Stage one requires oxidation of CH to create cyclohexanone (-One) and cyclohexanol (-Ol). This is done through the ketone and alcohol oil process (KA-oil) and operationalizes a cobalt or manganese catalyst at around 150°C and at 10-20 bar of air. Stage two derives AA from the KA-oil mixture by applying nitric acid as an oxidant [126]. The majority of AA is produced using this process. However, this process involves significant amount of recycling (more than 90%) of unreacted cyclohexane due to its low conversion (5–10%). The reaction is deliberately performed at such low conversion to obtain high selectivity (70–85%) of KA products. This recycling of ≥90% certainly incurs higher operation costs. Besides, there are also some additional environmental implications. Particularly, the usage of nitric acid in the second step results in the formation of NO\textsubscript{x} by-products, which can be held responsible for smog, acid rain, and ozone depletion. Therefore, there is a need to explore new possible routes for the production of AA. This is the real motivation to uncover a more environment-friendly route using nano gold catalysts of the present study. Scheme 3 depicts some other possible options for the production of AA. Although there are many potential options available, direct synthesis of AA from CH in a one-step process using O\textsubscript{2} is considered to be a more suitable option from both environment-friendly and economic points of view.

This research has also addressed oxidizing adipic acid from cyclohexane and the associated catalytic activity in addition to the oxidation of benzaldehyde from benzyl alcohol using supported AuNPs [126]. Section 5.1 outlines the type of catalysts used in this instance. The
reaction conditions applied for determining the influence of support in CH oxidation were: 10 ml CH, 20 ml solvent (acetonitrile), 0.3 g catalyst, 0.1 g TBHP, \( pO_2 = 10 \text{ bar} \), \( t = 4 \text{ h} \), 1500 rpm, \( T = 130^\circ \text{C} \). The reaction resulted in adipic acid (AA), cyclohexanone (-One) and cyclohexanol (-Ol). Combining ketone and alcohol products is commonly denoted as KA (as is the case with commercially producing AA from cyclohexanone and cyclohexanol). Succinic acid, CO, CO\(_2\), and glutaric acid are expected by-products, formed in small quantities. However, these reactions are intended to produce AA.

Figure 6 depicts the impact of the nature of oxide support on CH to AA oxidation activity. It is evident from the figure that the characteristics of the supports clearly influence both the product selectivity as well as CH conversion. It is also clear that AuNPs supported on TiO\(_2\) gave the best performance compared to other supports. Similarly, even in case of benzyl alcohol oxidation, the same Au/TiO\(_2\) nanocatalyst was found to exhibit superior performance (Figure 6). The best performance of Au/TiO\(_2\) catalyst is due to (i) the formation of small AuNPs...
(2–3 nm), (ii) high dispersion of Au, (iii) higher active metal (Au) area, and (iv) the high surface enrichment of Au species compared to other catalysts applied in this study [126]. This finding further confirms that catalyst performance is strongly dependent on the size of Au particles. As discussed earlier, the poor catalytic performance was observed over the MgO- and CaO-supported catalysts. In addition, these catalysts also produce high amounts of unwanted by-products (including CO and CO₂), and thus they should not be considered as appropriate supports for this reaction. Conversion of CH achieved over TiO₂ was 16.4%, and the selectivity of AA was 21.6%.

Moreover, we have also made attempts to find some correlations between the superior performances of AuNPs supported on TiO₂ compared to other types of catalysts. Such correlation between the Au particle size, surface gold-to-support atomic ratio (Au/SU) derived from XPS and catalytic performance is presented in Figure 7. It is clear from the figure that the activity results are in good agreement with the particle size of Au, and therefore the conversion of CH and selectivity to AA changed relatively in a similar way as that of Au size. Unsurprisingly, the smallest gold nanoparticles supported on TiO₂ showed the best catalytic activity. The conversion of CH decreased from 26% using Au/TiO₂ catalyst to 9% using Au/CaO. On the other hand, the Au size increased from 2 nm in case of using TiO₂ to 6–8 nm using CaO.
Moreover, the XPS results also confirmed that the surface region was indeed enriched in Au which hence the catalytic activity of gold catalyst. We can obviously observe from Figure 7 that the gold nanoparticles supported on TiO$_2$ showed the highest surface Au/SU atomic ratios (SU = different supports) compared to others types of supports, and hence enhanced performance as well. In addition, supporting evidence for the beneficial effect of TiO$_2$ can be the high dispersion of Au and high active Au metal area. On the whole, better performance of TiO$_2$ is certainly due to the presence of small Au particles, high Au enrichment in the near-surface-region, high dispersion, and higher active metal area.

![Figure 7](Image)

**Figure 7.** Correlation between cyclohexane conversion (X-CH), selectivity to adipic acid (S-AA) and to KA oil (S-KA), and catalyst properties of supported AuNPs.

5.3. Supported metal nanoparticles for dye's photodegradation

Metal nanoparticles (e.g., Au, Cu, and Ag) have recently begun to generate attention regarding their uses in electronics, optics, and the environment, and as catalysts [128–130]. One important green technology application is the use of combined M and metal oxide supports to remove dyes photocatalytically from water. Toward this end, we looked at the potential of using gold MNPs supported on TiO$_2$ and ZnO under UVA light irradiation to test their photocatalytic activity toward photodegradation of rhodamine B (RB) in aqueous medium [128]. The
procedure previously described was used to prepare the catalysts. The UV-vis absorption spectra of a 20 mg/L rhodamine B solution is demonstrated in Figure 8a and b, which shows the results before and after UVA light irradiation at 20-minute intervals and in the presence of one of two photocatalysts: Au/ZnO and Au/TiO₂. Even before introducing light irradiation, a test was deployed to determine if dye absorption transpired on the catalyst surface. According to the tests, there was a slight amount of absorption (as shown by a diminished RB concentration), which was comparable between the both photocatalysts applied. Peak absorption with Au/TiO₂ was observed to occur at 554 nm, and thereafter decreased during irradiation, until full elimination after 100 min (Figure 8a). Peak absorption with Au/ZnO occurred at RB 554 nm, and likewise decreased with exposure to irradiation, reaching full RB depletion at 80 min (Figure 8b). The former’s 100-minute degradation time was slower, indicating that Au/ZnO has superior photocatalytic activity.

![Figure 8. Absorption spectra of RB at different times after UV irradiation using Au/TiO₂ (a) and Au/ZnO (b) as AuNPs photocatalysts.](image-url)

Furthermore, the degradation plot depicted in Figure 9 shows further details, demonstrating the photocatalytic degradation of RB in aqueous medium using undoped and Au-doped photocatalysts, which is dependent on the nature of the support. For example, undoped TiO₂ and ZnO have similar RB photodegradation reactions in the first hour, with approximately 73% of the present RB dye present in the solution. Approximately 98% was achieved after 160 min, with constant efficiency following. It is possible that the photocatalytic performance was so stable beyond 160 min due to the accumulated degradation products at the photocatalyst surface. However, enhanced RB photodegradation performance was achieved by doping both TiO₂ and ZnO with 1 wt.% AuNPs. In this case, 87% degradation was achieved following 60 min with Au/TiO₂ and 96% with ZnO in the same time. Enhanced photocatalytic degrading capability could potentially be due to AuNPs’ electron buffering, which results in slower recombination of pairs of electrons and holes [131]. Degradation efficiency of Au/ZnO rose to 97% when reaction time was increased to 80 min, and remained stable at this stage. Au/TiO₂ saw the same results when reaction time was increased to 100 min. This plateau behavior is likely due to reaction product deposits on the surface of the photocatalysts.
6. Conclusions and outlook

Particles having the size from 1 to 100 nm are termed as nanoparticles, and in nanotechnology, small particles behave as an identical unit in both chemical and physical aspects. Supporting such metal nanoparticles is found to be a great tool where they can be used to develop different green catalytic applications. The interesting phenomenon about these materials is that their high surface area-to-volume ratio of solid-supported metal particles is essentially responsible for their catalytic properties. In this chapter, the recent reported methods for fabricating MNPs have been reviewed. However, the selection of such method is found to be dependent on the desired application. We have also shown that a deeper knowledge to understand the catalytic properties of MNPs is needed to explore the potential application of these MNPs. Therefore, different spectroscopic and microscopic techniques used to understand the catalytic parameters in MNPs have been briefly reviewed. A number of important green reactions using supported MNPs have been presented in this chapter. Quite recently, we have successfully reported that supported MNPs catalyst is an efficient catalyst for different applications such as the oxidation of cyclohexane to adipic acid. Here, one can clearly notice that the size of Au and the nature of support applied matter a lot. In this context, smaller Au particles revealed superior performance compared to their corresponding counterparts. Based on this, the Au/TiO$_2$ exhibited the best performance compared to other catalysts of the same series for both benzyl alcohol oxidation to benzaldehyde and cyclohexane oxidation to adipic acid. In addition, the application of supported MNPs as efficient photocatalysts was also reported.
Overall, recent advances have been made in governing metal particle morphology, size, and synthetic methodology. Prior art research demonstrates that synthetic approaches and adjustments to chemical and redox environments can result in control over the final form and morphology of nanoparticles. Using oxide carriers to synthesize stable MNPs has environment-friendly implications due to the fact that MNPs have unique properties compared to bulk or isolated atoms. When these MNPs interact successfully with the support, they may stabilize the surface and may generate Lewis sites that may catalyze certain catalytic reactions. However, the interaction degree depends upon the kind of support applied. Therefore, MNPs have varied and new environment-friendly applications. In view of this, highly unsaturated systems are considered superior as they provide larger number of active surface metal sites that may allow efficient interactions between the support and active phases. The ability to stabilize and control other MNPs (such as Pd, Rh, Pt, etc.) would be an excellent advancement, allowing concepts and reactions now possible with homogenous catalysis to be used in a heterogeneous context. However, the commercial application is limited by their activity and selectivity properties. Stability and durability of supported MNPs have implications for productivity. The research presented here was driven by determining the role of supported MNPs in various reactions taking place, either in liquid or gas phase, as well as their associated catalytic processes. Future research may address the creation of bifunctional or multifunctional MNPs, which may have implications for existing reactions or even new outputs. Furthermore, rational approaches need to be worked out in the coming years to sequence various functionalities in the direction of developing novel, attractive, and ecofriendly nanostructured metal nanoparticles for various catalytic applications.

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