Green Synthesis of Metal and Metal Oxide Nanoparticles: Principles of Green Chemistry and Raw Materials

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Abstract: Increased request for metal and metal oxide nanoparticles nanoparticles has led to their large-scale production using high-energy methods with various toxic solvents. This cause environmental contamination, thus eco-friendly “green” synthesis methods has become necessary. An alternative way to synthesize metal nanoparticles includes using bioresources, such as plants and plant products, bacteria, fungi, yeast, algae, etc. “Green” synthesis has low toxicity, is safe for human health and environment compared to other methods, meaning it is the best approach for obtaining metal and metal oxide nanoparticles. This review reveals 12 principles of “green” chemistry and examples of biological components suitable for “green” synthesis, as well as modern scientific research of eco-friendly synthesis methods of magnetic and metal nanoparticles. Particularly, using extracts of green tea, fruits, roots, leaves, etc., can be used for the „green” synthesis of spinel magnetic NPs. “Green” nanoparticles are being widely used as antimicrobials, photocatalysts and adsorbents. “Green” magnetic nanoparticles demonstrate low toxicity and high biocompatibility, which allows for their biomedical application, especially for targeted drug delivery, contrast imaging and magnetic hyperthermia applications. The synthesis of silver, gold, platinum and palladium nanoparticles using extracts from fungi, red algae, fruits, etc., has been described.

Keywords: green synthesis; magnetite; spinel ferrite; metal nanoparticles

1. Introduction

Nowadays, a new page is turning in the history of chemistry, connected with the development of a new integrated scientific direction—“green” chemistry. “Green” chemistry is interdisciplinary: there is an integration of synthetic organic chemistry with analytical chemistry, physical chemistry, toxicology, microbiology, biotechnology and engineering. The goal of “green” chemistry is to develop technologies for more efficient chemical reactions. “Green” chemistry aims to prevent pollution in the very early stages of the planning and implementation of chemical processes and covers all types and aspects of chemical processes to minimize the environmental risks. The problems within the competence of “green” chemistry can be categorized into two main areas. The first relates to the processing and utilization of environmentally hazardous waste and by-products of the chemical industry. The second, more promising, involves the development of new industrial processes to eliminate or minimize the formation and use of harmful products [1]. “Green” chemistry allows to obtain the necessary substance in the safest possible way. It provides the selection of raw materials and process schemes, which generally exclude the use of harmful...
substances, toxic and hazardous chemicals, and focuses on industrial processes that do not pollute the environment and lay the responsibility for the products on the scientists and manufacturers [2].

As a scientific field, “green” chemistry appeared in the United States in the 1990s. European countries have been implementing the most advanced laws on “green” technologies. In recent years, new reaction schemes and processes have been developed, designed to drastically reduce the burden of chemical production on the environment, to minimize the processing and utilization of hazardous substances and harmful by-products [1]. “Green” chemistry is already moving in three major directions: new ways of synthesis (using catalysts); replacement of traditional organic solvents (particularly, the use of supercritical CO₂); renewable source reagents (i.e., non-petroleum products) [3].

In “green” chemistry, fundamentally new constructs such as “ideal process”, “ideal product” and “ideal consumer” are used [4]. The ideal process is a simple, eco-friendly, one-stage process, effective at the molecular level, with the use of renewable raw materials, which provides maximum yield. The ideal product requires a minimum of energy and packaging, is safe, recyclable and fully degradable by microorganisms [1]. Usually the main focus is on the production process and the final product, and the consumer is absent in this scheme. In “green” chemistry, the image of the “ideal consumer” is present—he uses a minimum number of goods, understands the need to preserve the environment. New research frontiers and new terms have been introduced into “green” chemistry: “atom efficiency”, “innate safety”, “product life cycle analysis”, “ionic liquid”, “renewable energy”, “environmental efficiency”, “process intensification and integration”, etc.

This review is aimed to analysis of modern scientific research of eco-friendly “green” synthesis methods of magnetic and metal nanoparticles. “Green” nanoparticles are being widely used as antimicrobials, photocatalysts and adsorbents [5–7]. “Green” magnetic nanoparticles demonstrate low toxicity and high biocompatibility, which allows their biomedical application, especially for targeted drug delivery, contrast imaging and magnetic hyperthermia applications. The structure and morphology of synthesized magnetic nanoparticles can be characterized by scanning electron microscopy, transmission electron microscopy, energy-dispersive analysis, X-ray diffraction analysis, X-ray photoelectron spectroscopy, FTIR spectroscopy, Raman spectroscopy and magnetic force microscopy. Magnetic force microscopy (MFM) is a scatter-sensitive technique with a resolution of up to 10 nm that can detect weak magnetic fields. MFM is a universal method of analysis of magnetic nanoparticles due to the simple requirements for sample preparation, the ability to work in air, vacuum or liquid medium [8]. Due to this, MFM is a powerful tool for imaging of magnetic NPs, characterizing their size and morphology. For superparamagnetic nanoparticles, MFM has been especially useful to evaluate magnetic moment, magnetic anisotropy, magnetization curves and the effect of aggregation in particles [9]. The ability of MFM to detect superparamagnetic and low-coercive magnetic nanoparticles and the interpretation of the obtained MFM images are the subject of many research [10–15]. Torre et al. [10] demonstrated the ability of magnetic force microscopy (MFM) to quantify magnetic textures at room temperature. MFM measurements were performed for magnetic nanoparticles of iron oxide with a diameter of 11 nm. The obtained images of nanofilms, which were applied to the substrate, indicated linear magnetic chains of nanoparticles of several hundred nanometers. In a study [11] the authors quantified the magnetization of individual magnetic NPs using MFM. Cordova et al. [12] used MFM to analyze the superparamagnetic iron oxide NPs in air, liquid medium and inside thin polymer films. The authors of work [13] used MFM to analyze the magnetite NPs with different sizes (from 10 nm to 100 nm), which are embedded in polymer films with different thickness. Therefore, MFM allows to quantify the magnetic properties of both single nanoparticles and nanoparticles in non-magnetic matrices (e.g., polymers).
2. The Principles of “Green” Chemistry

In 1998 Paul Anastas and John Warner in their book “Green Chemistry: Theory and Practice”, Ref. [2] formulated 12 principles of “green” chemistry. They recommend the scientists, industrialists and government officials to direct their activities to reduce or eliminate the use of hazardous materials and chemical processes. These 12 principles, due to their relevance, usefulness and specificity, have made a significant contribution to the expansion and formation of a new philosophy.

The principles are following (Figure 1):

- prevention of waste (chemical synthesis design that prevents waste rather than its disposal or utilization);
- maximum increase of components—“atom economy” (design of synthesis to maximize raw materials ratio in the final product with the least or no amount of waste);
- development of less dangerous chemical syntheses (generating and using substances with minimal or zero toxicity);
- design of safe chemicals and products (chemicals that are effective yet non-toxic);
- use of safe solvents and reaction conditions (minimize or exclude the use of solvents or other auxiliary chemicals, and if necessary—use the safest of them);
- increase energy efficiency (identify and minimize the consequences caused by using energy in chemical synthesis. Initiate chemical reactions at room temperature and pressure, if possible);
- use of renewable raw materials (sources of renewable raw materials are agricultural products or waste);
- avoidance of chemical derivatives (minimize or eliminate the use of blocking or protective groups or any temporary modifications, if possible);
- use of non-stoichiometric catalysts (minimization of waste by implementation of catalytic reactions, use of effective catalysts in small quantities that can promote the reaction repeatedly);
- design of degradable chemicals and products (non-persistent, which decompose into safe substances);
- real-time analysis of pollution (elimination or minimization of by-products through interfering with the process during synthesis);
- minimizing the possibility of accidents (such as releases, explosions and fires) through designing safer chemicals and their physical forms (solid, liquid or gaseous).

![Figure 1. Principles of “green” chemistry.](image-url)

R.A. Bourne et al. [3] presents 12 principles of “green” chemistry using the abbreviation “PRODUCTIVELY” (Figure 2). Let us look closely to each of the 12 principles [16,17].
2.1. Prevention (Reducing) of Waste/by-Products

The major principle is called the prevention principle, and the other principles are the “how-to’s” to achieve it. The best way is to carry out the synthesis with zero or no waste (by-products), because the costs associated with the waste disposal significantly increase the total cost of production. Even unreacted raw materials are part of the waste. Therefore, we should avoid the generation of waste (or by-products), which causes pollution when dumped into the atmosphere, sea or land and requires cleaning costs [16]. Introduction of the E-factor (“by-products/final product” ratio) by Roger Sheldon of Delft University (the Netherlands) was an important innovation of “green” chemistry. It characterizes the loss per 1 kg of target product, allows to compare chemical production technologies, and is crucial for attracting attention of global chemical and pharmaceutical industries to the issue of waste [18].

2.2. Maximum Inclusion of Reagents (Source Materials) in the Final Product

When one mole of reagent results in one mole of product, the yield is 100%. Chemists around the world consider the reaction quite effective, when the yield is about 90%. However, product yield calculations can create excessive waste (or by-products). Typical examples, such as the Grignard or Wittig reactions, confirm those statement. Aforementioned reactions can result in 100% yield, but ignore the number of by-products. The reaction is considered “green” if there is a maximal inclusion of precursors in the final product [16].

The term “atom economy” was introduced in 1973 and has become a basic concept among researchers in this field of chemistry. The main goal of atom economy was to overpass the limitations of traditional “profitability”, the amount of final products used in calculating the effectiveness of reactions. For example, to calculate yields, chemists considered the effectiveness and amount of just the basic chemical product they chose (“target molecules”), excluding possibly hazardous by-products. The atom economy takes into account all the components and reagents of the reactions, thus providing a reliable indicator of whether pollutants are formed during the reaction. Green chemistry has proven the reduction of pollution to be possible through atom economy, which relies on such processes as hydrogenation, metathesis and cycloaddition.

![Figure 2. The principles of “green” chemistry.](image-url)
2.3. Prevention or Minimization of Harmful Products

“Green” chemistry’s major principle is to avoid or reduce the formation of hazardous products. The danger to workers can be decreased by using protective clothing, respirators, etc. This, however, increases the cost of production. To avoid risks, “green” chemistry has found a scientific solution to such situations [16].

2.4. Development of Safer Chemicals

A priority is to ensure that the synthesized chemicals (dyes, paints, adhesives, cosmetics, pharmaceuticals, etc.) are harmless. An example of a dangerous substance is thalidomide (introduced in 1961) for nausea and vomiting of pregnancy. Has been proven that the children born to women taking this drug had different birth defects (including missing or deformed limbs). Subsequently, thalidomide was banned and strict rules for testing new drugs were implemented. With the development of technology, it has become possible to produce safer chemicals by manipulating the molecular structure of substances [16].

2.5. Energy Requirements for the Chemical Synthesis

The minimum energy requirement must be adhered to any chemical processes. For instance, if the precursors are soluble in a specific solvent, the reaction mixture must be heated for some time or until completion. In such circumstance, the time required to complete the reaction should be minimal with a minimum amount of energy required. A catalyst can be used to reduce the energy needs of the reaction. In case of an exothermic reaction, large cooling is sometimes required. Sometimes the final product must be purified by ultrafiltration, distillation, or recrystallization. All these stages are energy consuming and increase the total cost. Final energy requirements can be minimal if the process is planned so that there is no need for separation or purification [16].

2.6. Selection of Proper Solvent

The chosen solvent should not lead to any environmental contaminations or health hazards. The use of liquid or supercritical CO$_2$ should be studied. If possible, the reaction should be performed in the water medium or without a solvent. The best method is to perform the reaction in the solid phase. One of the main problems with many solvents is their volatility, which can be harmful to the health and ecosystems. In order to avoid this, immobilized solvents can be used. They maintain the solubility of the material, are non-volatile and safe. Immobilization can be carried out by attaching the solid substances to a solid phase or by binding the solvent molecule directly to the polymer matrix. Several newly discovered polymeric substances as solvents have been found to be non-hazardous [16].

2.7. Selection of Proper Source Materials

Source materials are derived from renewable or non-renewable materials. Petrochemicals are usually derived from crude oil, that is not a renewable source. Raw materials obtained from agricultural or organic products are called renewable. However, such factors as crop failure, etc., may interfere with constant supply of agricultural products. Substances such as carbon dioxide (formed naturally or synthetically) and methane (derived from natural sources) are in sufficient quantities. They are considered as renewable raw sources [16].

2.8. The Use of Catalysts

Catalysts promote the reaction without being consumed and included in the final product. Thus, they should be used whenever possible. The benefits of catalysts include: (i) better product yield; (ii) the reaction becomes possible in cases where it does not occur normally; (iii) increased selectivity. In addition, the use of catalysts has significant advantages in energy demand, better utilization of raw materials and waste minimization. With advances in the catalysts selectivity, certain “green” synthesis reactions have become very convenient [16].
2.9. Biodegradation of Obtained Products

The problem of non-biodegradable products is especially common with insecticides and polymers. Farmers are using different types of insecticides in order to protect crops from insects. Widely used insecticides include less stable (carbamates, organophosphates) and more stable (chlorinated hydrocarbons). Although the latter are certainly effective, they are usually bioaccumulated in flora and fauna and included in the food chain. The insecticides cause a decrease in the population of beneficial insects and animals (honey bees, butterflies, mites, etc.). Given the above, it is crucial that any synthesized product is biodegradable and non-toxic [16].

2.10. Strengthening Analytical Methods for Controlling Harmful Compounds

Analytical methods should be designed to require minimal use of chemicals. For example, processing some unreacted chemicals to complete the reaction. It is also useful to place sensors to track the formation of toxic by-products during a chemical process [16].

2.11. Development of Production Units

The importance of accident prevention in production units cannot be overstated. A number of industrial accidents have occurred and caused not only in the loss of thousands of lives but also in lifelong disabilities. Production facilities should be fabricated to exclude the accidents possibility caused by toxicity, explosions, fires, etc., during operation [16]. Many industrial enterprises have welcomed the proposed 12 principles and have made some progress in improving the safety of their chemical plants. For example, the world-famous company Pfizer has developed a new technology for the production of sildenafil citrate. While the old technology required 1300 L of solvent containing chlorine, the new—only 6.5 L of safe solvent. As a result, the mentioned E-factor of such production decreased from 105 to 6, and the pharmaceutical giant itself received a prize from the British government [3].

3. Raw Materials for “Green” Chemistry

3.1. Transition to Renewable Raw Materials

During last 80 years, the chemical industry has been based on natural gas and crude oil as the main raw sources. However, today there is a trend towards transition from fossil to renewable raw resources, such as carbohydrates and biomass-derived triglycerides [18]. A partial transition to renewable energy sources is desirable for such reasons as biocompatibility, biodegradability and lesser toxicity. Products based on renewable raw materials are obtained from carbon dioxide and water via photosynthesis and after being used, are eventually returned to the biosphere as CO$_2$ and H$_2$O through biodegradation. They are becoming more reliable and cheaper compared to the rapidly increasing gas and oil prices. The developing of “green” products, which can replace petroleum-based products and the implementing of «green synthesis» methods for the chemicals production from biomass, are the main in the transition to renewable raw materials. For example, the catalysts (like modified corn starch with surface -SO$_3$H/-NH$_2$ groups) for chemical reactions can be obtained from biomass [18,19].

3.2. Biological Components for “Green” Synthesis

In order to obtain “green” nanoparticles with required shape, size, and properties, two synthesis principles are being considered: “top-down” and “bottom-up” [20]. In the “bottom-up” approach, the NPs are formed first, and then assembled into the final material. The benefit of the „bottom-up“ principle is the opportunity of obtaining small metal NPs with uniform chemical composition. In the „top-down“ approach, the source material is reduced in size by physical (e.g., mechanical) or chemical methods. The main disadvantage of this approach is the defects of the material surface, which can significantly affect the properties of metal nanoparticles [21].
The environmentally friendly and sustainable synthesis methods are using in “green” chemistry in order to prevent the formation of toxic by-products [22–24]. Different biological materials, such as plant extracts, algae, fungi, bacteria, etc., are used in the “green” synthesis of metal and metal oxide NPs [25–36]. The use of plant extracts is fairly easy for obtaining NPs on a large scale, compared to the synthesis with fungi and bacteria [20,37,38]. Various factors (pH, pressure, temperature, solvent type) affect the “green” synthesis techniques. However, the key role is belongs to the phytochemicals, which are presented in plant extracts (roots, leaves, stems, fruits): ascorbic acids, phenols, carboxylic acids, terpenoids, amides, flavones, aldehydes, ketones, etc. [39–42]. These components reduce metal salts to metal NPs [20]. There are different mechanisms of NPs formation using microorganisms [38]. A huge variety of nature biological materials, including plants [43–51], algae [52–56], fungi [57–61], yeast [62–65], bacteria [66–69], viruses [70], etc., can be used for the synthesis of “green” NPs (Figure 3).

![Figure 3. Various natural resources used for the synthesis of “green” nanoparticles.](image)

### 3.2.1. Bacteria

Bacteria are widely used for genetic engineering, bioremediation and bioextraction. Various types of bacteria are able to reduce metal ions and are important in NPs obtaining. In particular, prokaryotic bacteria and actinomycetes are widely used for the synthesis of metal or metal oxide NPs [20]. Some examples of bacterial strains that are widely used for the obtaining of bio-reduced silver NPs with different size and morphologies include: Shewanella oneidensis, Arthrobacter gangotriensis, Enterobacter cloaceae, Bacillus cereus, Bacillus subtilis, Bacillus subtilis, Bacillus megaterium D01 [21].

### 3.2.2. Fungi

Fungus-mediated metal/metal oxide NPs biosynthesis is also very productive for obtaining monodisperse NPs with desired morphologies. They are preferable for the production of metal and metal oxide NPs because of various intracellular enzymes. Proper
fungi can synthesize more nanoparticles than bacteria. Furthermore, the presence of reducing components, enzymes, proteins in their cells grants them advantage over other organisms. The probable mechanism of metal nanoparticle formation is an enzymatic reduction (reductase) in the cell wall or inside the fungal cell [20]. A clear advantage of fungi in the synthesis of nanoparticles is the simplicity of their scaling (e.g., using the method of thin solid substrate fermentation). The fungi are highly effective secretors of extracellular enzymes. Thus, it is easy to obtain large-scale production of enzymes. More advantages of using fungi in the “green” synthesis of metal nanoparticles include economic viability and easy biomass handling. However, there is a significant disadvantage of using these bioformations in the synthesis of nanoparticles as the genetic manipulation of eukaryotes is much more complex than prokaryotes [21].

3.2.3. Yeast

Yeast is a unicellular microorganism that is present in eukaryotic cells. Only 1500 species of yeast have been identified. Numerous research groups have reported the successful synthesis of nanoparticles/nanomaterials using yeast. Many different types are used to produce countless metal nanoparticles [20,62].

3.2.4. Plants

Plants can accumulate heavy metals in leaves, roots, fruits, etc. Therefore, synthesis using plant extracts attract attention as simple, effective, cheap and feasible methods for obtaining nanoparticles [71–73]. Various plants can be used for reduction and stabilization of metal nanoparticles during synthesis. Many researchers use “green” synthesis to obtain metal oxide NPs using plant extracts for various applications [74–78]. The plant extracts are mixed with the solutions of metal precursors under different reaction conditions [79–82]. Such parameters as temperature, pH, metal salt concentration, types and concentration of phytochemicals affect the stability and the rate of NPs formation. Biologically active compounds found in plants (Figure 4) due to the presence of functional groups are able to reduce metal ions much faster than bacteria or fungi. Amides, carboxylic acids, aldehydes, ketones, sugars, terpenoids and flavones are among essential phytochemicals, which are responsible for the NPs bioreduction [83–86].

![Figure 4. Plant phytochemicals.](image-url)
Plants contain biologically active compounds (carbohydrates, coenzymes and proteins) with excellent ability for the reduction of metal salts to NPs [87,88]. The syntheses of gold [89–94] and silver [95–101] nanoparticles involving plant extracts were the first to be studied. Various plants are used: lemon grass (Cymbopogon flexuosus), mustard (Brassica juncea), coriander (Coriandrum sativum), grape (Vitis), Ginkgo Biloba, Cydonia oblonga, neem (Azadirachta indica), lemon (Citrus limon), tulsi (Ocimum sanctum), oats (Avena sativa) and aloe vera (Aloe barbadensis). The Zn, Ni, Co and Cu NPs is obtained using sunflower (Helianthus annuus), alfalfa (Medicago sativa) and mustard (Brassica juncea). ZnO NPs were also obtained from a wide number of plant extracts, such as green tea (Camellia sinensis), China rose (Hibiscus rosa-sinensis), copperleaf (Acalypha indica), coriander (Coriandrum sativum) and crown flower (Calotropis gigantea) [20,102–104].

3.3. Solvent-Based “Green” Synthesis

Solvent systems are the main components in the synthesis process. Most common solvents are harmful. One of the well-known solvents is benzene, which causes cancer in humans. Some of the aromatic solvents, such as toluene, can cause brain damage, affect speech, vision, and cause problems with kidney. Halogen-containing solvents, such as dichloromethane, carbon tetrachloride, perchloroethylene and chloroform, are also commonly used and having been recognized as carcinogens [16].

Water is an ideal solvent, as it is the cheapest and most available, and has been used in the synthesis of various nanoparticles since the beginning of nanoscience and nanotechnology [20]. Carbon dioxide is a universal solvent, which is used as liquid CO$_2$ or supercritical CO$_2$. The gas usually turns into a liquid state after increased pressure. However, if the CO$_2$ is put at a temperature above 31 °C and at a supercritical pressure equal 7.38 kPa, a supercritical liquid is formed [16]. Supercritical and ionic liquids are some of the best solvents applied in „green“ synthesis. Ionic liquids consist of ions with melting points below 100 °C. They are called „ionic liquids at room temperature“. The ability of ionic liquids to be both reducing and protective agents simplifies the process of nanoparticle synthesis. Ionic liquids can be hydrophilic or hydrophobic based on the anions and cations nature [20]. The advantages of using ionic liquids as solvents are: (i) easy solubility of many organic compounds, metals and gases in ionic liquids; (ii) constructive thermal stability when operating in a wide temperature range (ionic liquids have a 3–4 times larger temperature range of synthesis, compared with water); (iii) ionic liquids do not coordinate, compared to other polar solvents or alcohols; (iv) ionic liquids do not evaporate as volatile organic solvents; (v) ionic liquids are amphilites due to the presence of cations and anions [20]. However, the main disadvantage of ionic liquids is their non-biodegradability. Thus the new ionic liquids with high biodegradation efficiency are being developed.

4. “Green” Synthesis and the Use of Magnetic Nanoparticles

4.1. Effect of Magnetic Nanoparticles on Environmental Restoration

In recent years, the environment has been polluted due to the excessive use of fertilizers and pesticides [105–108]. Extensive research work is underway to study various aspects of environmental restoration using magnetic nanoparticles, which includes cleaning the atmosphere, soil, sedimentary rocks, groundwater and surface water [109]. The synthesis of magnetic nanoparticles is the subject of number of systematic research related to their technological applications [110]. Magnetic nanoparticles have demonstrate high potential in environmental and biomedical applications [109,111,112]. Magnetic NPs are able to detoxify the environment due to the high specific surface area. They act as a “superabsorbent” for many contaminants, converting them into non-toxic forms. Magnetic nanoparticles are usually consists from Fe, Ni, Co or their oxides, such as magnetite (Fe$_3$O$_4$), maghemite (γ-Fe$_2$O$_3$), cobalt ferrite (CoFe$_2$O$_4$) etc. They can be controlled by magnetic fields [109]. The earliest reports of the synthesis of magnetic iron oxide particles, which form relatively stable colloids, date back to the 1930s. The first stable suspension of magnetic particles was obtained in 1965 by Steve Papell. His magnetic fluid was a dispersion of
crushed particles of magnetite (Fe₃O₄) (diameter <25 mm) modified with oleic acid. These particles were dispersed in non-polar solvents (carrier), forming a stable magnetic fluid, which was used to give the fuel its magnetic properties. Rosensweig obtained several types of magnetic fluids based on the dispersion of crushed Fe₃O₄ particles in various carriers, such as kerosene, water, fluorocarbons and esters. In 1982, Massart obtained magnetic fluids by chemical means, which involved the co-precipitation of Fe(II) and Fe(III) hydroxides, and modified the co-precipitation method to obtain ultrastable and highly concentrated magnetic fluids with different magnetic particles based on spinel ferrites, such as (M₁₋ₓ²⁺Feₓ³⁺)ₐ[Fe₂₋ₓ³⁺Mₓ²⁺]ₘO₄ (M = Mn, Co, Ni, Cu, Zn). The stability of such dispersions is achieved via hydrothermal treatment of samples with Fe(NO₃)₃, which leads to the formation of a protective layer rich in iron, which passivates the nanoparticle surface [110]. However, these techniques do not allow good control of the nanoparticles morphology. Therefore, the researchers’ efforts were focused on finding ways of synthesis where better control of the size and shape of magnetic nanoparticles can be achieved. Many scientific studies present different concepts for controlling the morphology of synthesized magnetic nanoparticles, but none of them fully meet all the principles of “green” chemistry [110]. There are several types of magnetic nanoparticles: ferrites, core-shell NPs (magnetic core plus shell from SiO₂, for example), metal NPs (are pyrophoric and react to oxidants, what complicates their use and causes undesired side effects) etc. [113]. Magnetic NPs, coated with SiO₂, have a few advantages over metal nanoparticles: higher chemical stability; narrow size distribution; higher colloidal stability; adjustable magnetic moment by the size of the nanoparticle cluster; preserved superparamagnetic properties; the SiO₂ surface allows direct covalent functionalization [113].

4.2. Superparamagnetic Iron Oxide (Magnetite) Nanoparticles and Their Application

Magnetite Fe₃O₄ is a most common natural iron oxide with inverse spinel structure. The ferrous (Fe²⁺) ions occupy half of the octahedral positions of Fe₃O₄ spinel structure due to the higher field stabilization energy of black crystalline substances, and Fe³⁺ occupy the remaining octahedral positions and all tetrahedral positions. The magnetic behavior of Fe₃O₄ nanoparticles strongly relies on the synthesis method. In addition, the NPs morphology are key to magnetic properties of magnetite. Therefore, for better application, it is necessary to determine the optimal parameters of Fe₃O₄ nanoparticles [114].

Superparamagnetic Fe₃O₄ NPs are well known due to their unique properties: biodegradability, biocompatibility and non-toxic effect. If the nanoparticle size less than 50 nm in diameter, they are suitable for effective endocytosis with the use of drugs. Therefore, many synthesis methods (co-precipitation method, sol-gel method, hydrothermal synthesis, solid-state synthesis, flame spraying synthesis, thermal decomposition, solvothermal synthesis, etc.) to obtain Fe₃O₄ nanoparticles with the desired properties have been reported [114]. Figure 5 shows the possible use of Fe₃O₄ nanoparticles, namely: as a catalyst, for water purification (removal of heavy metal ions), lithium-ion batteries, biomedical applications, tissue engineering etc. All these studies show promising results and provide a platform for Fe₃O₄ nanoparticles, whose unique features provide great potential for their widespread use.
Maghemite (γ-Fe$_2$O$_3$) is isostructural with magnetite, but it has cation vacancies and quite similar general properties, even though maghemite is generally less magnetic but more stable than magnetite. It can be obtained by direct oxidation of magnetite. Although maghemite does not always need a shell for stabilization or a graft to the surface with polymers (such as polyethylene glycol), they are often used in the biological system and increase the half-life of nanoparticles in the blood. The increase in half-life is usually associated with a delay in the opsonization process, in which particles are targeted by phagocytic immune cells, resulting in rapid clearance to the liver or spleen through the reticuloendothelial system. Another way to increase the half-life is to add a biomolecular corona that interacts with biological systems. This corona can be the main element of the biological identity of the nanoparticle [115].

The “green” synthesis of magnetic nanoparticles uses a “bottom-up” approach, when metal atoms gather and form clusters, and then NPs [116–124]. Biologically active substances, which are presented in “green” sources, can reduce and stabilize nanoparticles during synthesis. This allows to control their shape and morphology required for specific applications [114,125–129].

4.3. “Green” Synthesis of Magnetic Fe$_3$O$_4$ Nanoparticles

Here are some examples of using “green” synthesis to obtain magnetic Fe$_3$O$_4$ nanoparticles, and their practical application (Figure 6). Venkateswarlu et al. [130] reported a removal of Pb(II) using DMSA-modified Fe$_3$O$_4$ (DMSA = dimercaptosuccinic acid). Fe$_3$O$_4$ were obtained by “green” method using Punica Granatum peel extract from FeCl$_3$·6H$_2$O and CH$_3$COONa precursors. Then, 0.926 g of dried product (Fe$_3$O$_4$) and 0.7288 g of DMSA were added to 40 mL of double distilled water, mixed together with ultrasound for 10 h at room temperature, and the pH was adjusted to 8 by adding 0.01M NaOH solution dropwise. After 10-h reaction, the obtained DMSA@Fe$_3$O$_4$ were separated under an external magnetic field, washed and dried at 90 °C in vacuum. These magnetic DMSA@Fe$_3$O$_4$ nanorods were used to remove Pb(II) from water medium. The adsorption capacity was equal to 46.18 mg/g at a dosage of 0.1 g/L and T = 301 K. The experimental data corresponded to the kinetic model of the pseudo-second order. Figure 6a shows the magnetic properties DMSA@Fe$_3$O$_4$ magnetic nanorods (MNRs) and Figure 6b shows TEM image of Fe$_3$O$_4$ and DMSA@Fe$_3$O$_4$ MNRs. The results indicate that the synthesized biogenic DMSA@Fe$_3$O$_4$ nanorods could be used as promised adsorbent for the Pb(II) removal.
Niraimathe et al. [131] obtained iron oxide NPs using iron sulfate (FeSO₄·7H₂O) and an aqueous extract of *Mimosa pudica* root containing mimosin, which acts as a reductant. IR spectroscopy approved the presence of biologically active plant molecules on the iron oxide NPs surfaces. The obtained iron oxide NPs can be used directly in the targeted delivery of pharmaceuticals.

Lunge et al. [132] successfully synthesized magnetic Fe₃O₄ NPs (MION-Tea) using tea waste as green reductant. MION-Tea exhibited supermagnetic nature (Figure 6c) and reveals the cuboidal/pyramidal shaped structure of Fe₃O₄ (magnetite) crystals. TEM analysis of MION-Tea shows that the formed NPs are in the range of 5–25 nm (Figure 6d). MION-Tea NPs were investigated as adsorbents for As(III) and As(V) removal from water medium. The adsorption capacity was 186.69 mg/g for As(III) and 153.8 mg/g for As(V). Comparison with known adsorbents revealed that MION-Tea has potential for the As(III) and As(V) ions removal.

Bahadur et al. [133] performed a detailed examination of the optical, thermal, magnetic and dielectric properties of citric acid modified superparamagnetic Fe₃O₄ NPs (Cit-USPMNs), obtained via a “green” co-precipitation route. Lemon juice was used as a source of citric acid. Cit-USPMNs were superparamagnetic with low coercivity and saturation magnetization from 31.4 emu/g to 61.8 emu/g (Figure 6e). The size of Cit-USPMNs was in the range of 11–15 nm (Figure 6f).

**Figure 6.** (a) VSM of DMSA@Fe₃O₄ MNRs. Upper inset shows DMSA@Fe₃O₄ MNRs dispersed in water and its magnetic separation and lower inset shows the enlargement of the hysteresis loop at low magnetic field (Adapted with permission from [130], Elsevier, 2019); (b) TEM image pattern of DMSA@Fe₃O₄ MNRs (Adapted with permission from [130], Elsevier, 2019); (c) VSM Magnetization curve of MION-Tea (insight nanoparticles attracted by magnetic retriever) (Adapted with permission from [132], Elsevier, 2014); (d) TEM image of MION-Tea (Adapted with permission from [132], Elsevier, 2014); (e) M-H loop for Cit-USPMNs (11nm) at room temperature (Adapted with permission from [133], Elsevier, 2017); (f) TEM image of Cit-USPMNs (11nm) (Adapted with permission from [133], Elsevier, 2017).

Kanagasubbulakshmi et al. [134] revealed that the *Lagenaria siceraria* leaf extract is suitable for “green” synthesis of magnetite nanoparticles with enhanced antimicrobial properties. The synthesized Fe₃O₄-NPs were of cubic shape and the size range from 30 to 100 nm. Phytochemicals, which are presented in the leaves, act as reducing agents.
The –OH and –COOH functional groups present in nanoparticles make them hydrophilic, so they do not require additional functional modification. The antimicrobial properties of the “green” Fe$_3$O$_4$-NPs were investigated against gram-negative (E. coli) and gram-positive (S. aureus) bacteria strains. It was concluded that “green” Fe$_3$O$_4$-NPs demonstrate great potential for biomedical applications.

Padhi et al. [135] reported a new single-stage hydrothermal synthesis of photocatalytically stable and magnetically separated g-Fe$_3$O$_4$/2RGO nanocomposite in the presence of Averrhoa carambola leaf extract (as a natural surfactant for multipurpose use) for water purification. The adopted hydrothermal process leads to good incorporation of g-Fe$_3$O$_4$ nanoparticles with an average size of $22 \pm 2$ nm into 2D sheets of graphene oxide (RGO). Averrhoa carambola leaf extract was crucial in modifying the structural, optical and electronic properties of Fe$_3$O$_4$ nanoparticles. At room temperature, the g-Fe$_3$O$_4$/2RGO nanocomposite showed 97% in Cr(VI) reduction (50 mg/L in 1 h) and 76% in phenol degradation (10 mg/L in 2 h) under visible light. A higher activity of g-Fe$_3$O$_4$/2RGO is due to the presence of RGO in situ, which led to better separation of photoexcited charge carriers ($e^-/h^+$). In addition, g-Fe$_3$O$_4$/2RGO nanocomposite exhibited better antimicrobial activity against three bacterial pathogens, such as Staphylococcus aureous (MTCC-737), Bacillus subtilis (MTCC-736) and Escherichia Coli (MTCC-443), compared to GO and standard antibiotics (30 µg). The study proves g-Fe$_3$O$_4$/2RGO nanocomposite to be potentially useful as a good antibacterial agent.

Kataria et al. [136] synthesized new biogenic “green” magnetic iron oxide NPs, loaded with sawdust carbon (SC) and functionalized with EDTA (EDTA@Fe$_3$O$_4$/SC), to remove Cd(II) from the aqueous medium. The adsorption capacity toward Cd(II) ions was 63.3 mg/g. The results of regeneration studies proved the modified EDTA@Fe$_3$O$_4$/SC to be promising, cheap and eco-friendly for the adsorption of heavy metals from water environment.

Ahmadian-Fard-Fini et al. [137] prepared the Fe$_3$O$_4$/carbon dots nanocomposite for E.coli bacteria detection. Carbon dots (CDs) were obtained via hydrothermal method using extracts of grapes, lemon and turmeric in the presence of ethylenediamine. Next, Fe$_3$O$_4$ (magnetite) nanoparticles were obtained using these biocompatible retaining reagents. Figure 7a shows VSM curve of Fe$_3$O$_4$-carbon dot nanocomposite, and Figure 7b shows TEM image of magnetite-carbon dot core-shell nanocomposite. The results reveal the quenching of the photoluminescence of nanocomposites by increasing the number of bacteria.

Nnadozie et al. [138] reported a “green” biosynthetic co-precipitation of magnetite nanoparticles using Chromolaena odorata root extract, which acted as a precipitator and binder for nanoparticles. The extracted phenolic plant component was alkalized with 28% aqueous ammonia solution to pH 13, and added while stirring into the precursor solution (Fe$_2^+$ and Fe$_3^+$ ions in a 1:2 molar ratio) at 6 mL/min rate. The experiment was performed under a steady stream of nitrogen to avoid oxidation. The particle sizes were in the range of 5.6–16.8 nm. The peaks of the particles’ absorption bands at 205 nm and 291 nm are attributed to the oscillations of the surface plasmons, and the calculated band gap of the particles is 1.97 eV. Based on the extract, 30-fold single-phase magnetite nanoparticles are formed with a reduced band gap compared to the raw Fe$_3$O$_4$ nanoparticles. The obtained NPs demonstrate good water-disperse and hydrophilic properties.

Khatami et al. [139] synthesized superparamagnetic iron oxide nanoparticles (SPIONs) using a zero-calorie natural sweetener (Stevia) for reduction and stabilization. SPIONs (less than 25 nm) were very stable due to the biomolecular coating, as the zeta potential (−41.1 mV) creates opposite forces between the nanoparticles and prevents them from assembling. Biogenic SPIONs were able to counteract the effects of oxidative metabolites, according to a study of antioxidant activity. Figure 7c shows VSM magnetization curves of iron oxide nanoparticles. The FE-SEM image of the synthesized SPIONs is shown in Figure 7d. Great magnetic and catalytic properties, biocompatibility and low toxicity prove their potential for biomedical applications.
Figure 7. (a) VSM curve and (b) TEM image of Fe3O4-CDs nanocomposite obtained using lemon extract (Adapted with permission from [137], Elsevier, 2018); (c) The VSM magnetization curves and (d) FE-SEM image of iron oxide nanoparticles, synthesized using a zero-calorie natural sweetener (Stevia) (Adapted with permission from [139], Elsevier, 2019).

The food industry expresses great interest in β-glucosidase enzyme because of its role in the transformation of food to obtain functional foods. Moradi et al. [140] covalently immobilized β-glucosidase on aminotanic acid modified with magnetic Fe3O4 nanoparticles (ATA-Fe3O4 MNPs) as a biocompatible nanoplatform with a modified polyaldehyde pullulan to increase the ability and strength of the nanoparticle to bind the enzyme. The highest percentage of loading and immobilization yield was obtained with a solution of 0.1 mg of enzyme per 1 mL of citrate buffer (pH = 6; 1 M), a solution of citrate buffer carrier (ATA-Fe3O4)—10 mg/3 mL (pH = 6; 1 M) and a solution of polyaldehyde pullulan—20% of the total volume of the reaction system. Optimal pH and temperature values were found for free enzyme (pH = 5; 30 °C) and immobilized enzyme (pH = 6; 40 °C). The immobilized β-glucosidase enzyme retains its activity up to 83% after 10 cycles, so its immobilization by this method is an effective way for improving the properties of the enzyme. Magnetic hysteresis loops of the Fe3O4 MNPs, ATA-Fe3O4 MNPs, BGL-ATA-Fe3O4 MNPs are shown in Figure 8a. Figure 8b shows TEM image of the ATA-Fe3O4 MNPs.

Karade et al. [141] received magnetic Fe3O4 nanoparticles (with particle sizes from ~20 to 25 nm) via modified “green” synthesis method using green tea extract as a reductant and ethylene glycol as a solvent. As observed, the reaction time strongly affected the magnetic and structural properties of magnetic nanoparticles. As time increased, the crystallite size also increased from 7.5 to 12 nm with an improvement in saturation magnetization (Figure 8c,d). Magnetic measurements revealed that nanoparticles were superparamagnetic at room temperature, ferromagnetic and superparamagnetic—at 60 K. Magnetite magnetic nanoparticles synthesized using green tea extract are promised for bioapplication due to their biocompatibility and high magnetization.

Fatimah et al. [142] used Parkia speciosa husk extract for the synthesis of magnetic nanoparticles. The obtained NPs consisted from magnetite and hematite particles with sizes in the range of 10–80 nm. The magnetic NPs exhibited an excellent photocatalytic
properties in the degradation of the bromophenol blue dye under UV and visible light. The synthesized “green” nanoparticles are promising as photocatalysts in the degradation of dyes from wastewater.

Figure 8. (a) Magnetic hysteresis loops of the Fe$_3$O$_4$ MNPs, ATA-Fe$_3$O$_4$ MNPs, BGL- ATA-Fe$_3$O$_4$ MNPs and (b) TEM image of the ATA-Fe$_3$O$_4$ MNPs (Adapted with permission from [140], Elsevier, 2019); (c) Change in crystallite size as a function of reaction time and (d) FE-SEM image of Fe$_3$O$_4$ MNPs synthesized using green tea extract (Adapted with permission from [141], Elsevier, 2018).

4.4. “Green” Synthesis of Spinel Magnetic Nanoparticles

Spinel ferrites with the general formula AB$_2$O$_4$ have great chemical, catalytic, adsorption and magnetic properties [143–148]. Spinel ferrites have attracted much attention due to their thermal and chemical resistance [149–154]. Nickel ferrite (NiFe$_2$O$_4$) is a major representative of spinel ferrites with the inverse spinel structure, which demonstrates ferromagnetism, originating from the magnetic moment of the antiparallel spins of metal ions (Ni$^{2+}$ and Fe$^{3+}$) [155]. Udhaya et al. [156] developed a simple auto-combustion method using albumen for the synthesis of nanocrystalline nickel ferrite (NiFe$_2$O$_4$). Egg white (albumen), which is used in the “green” synthesis, plays the role of fuel in the process of auto-combustion. The results of powder analysis and IR spectroscopy indicated that the synthesized nanoparticles are single-phase and the spinel structure is cubic with a particle size of 23 to 47 nm. The dielectric properties of the nickel ferrite were measured for different frequencies from 100 Hz to 1 MHz. It was concluded that the alternating conductivity increases with increasing frequency.

Al-Hunaiti et al. [157] developed a “green” synthesis of magnetic CuFe$_2$O$_4$ NPs with an average size of 20 nm using Azadirachta indica extract. Cupper ferrite NPs were tested as an effective catalyst for the aryalkanes oxidation without solvents, especially in direct oxidation of toluene to obtain the desired benzoic acid in mild and eco-friendly conditions.

Routray et al. [158] synthesized nanosized CoFe$_2$O$_4$ via automatic combustion method using Aloe vera and the solutions of precursors Fe(NO$_3$)$_3$·9H$_2$O and Co(NO$_3$)$_2$·6H$_2$O. FE-SEM microphotographs revealed the formation of a bud-like structure and the resulting particle size was approximately 50–65 nm. Magnetic properties, especially saturation magnetization, remanence magnetization and coercivity, were examined from the M-H
loops: 72.23 emu/g, 31.29 emu/g, 1519 Oe, respectively. Furthermore, a massive dielectric constant, low dielectric loss and variable conductivity of CoFe₂O₄ nanoparticles depended on the frequency (100 Hz–1 MHz), preparation method and grain size.

Madhukara Naik et al. [151] prepared spinel zinc ferrite (ZnFe₂O₄) nanoparticles using the juice of Limonia acidissima (wood-apple). ZnFe₂O₄ nanoparticles were obtained by adding zinc nitrate and ferric nitrate to 5 mL of Limonia acidissima juice (reducing agent). The proposed method led to obtain ZnFe₂O₄ nanoparticles with an average crystallite size of 20 nm (Figure 9a). The study of magnetic properties reveals a high saturation magnetization of ZnFe₂O₄. ZnFe₂O₄ nanoparticles exhibited effective photodegradation of Evans blue and methylene blue dyes when exposed to visible light. Furthermore, the antibacterial activity (Figure 9b) of the nanoparticles was investigated against both gram-negative and gram-positive bacteria.

Ciprofloxacin (CIP) is an antibiotic that is widely used to treat infections. It is mostly excreted in non-metabolized form and enters the water via wastewater discharge. The aim of research by Malakootian et al. [159] was to synthesize ZnFe₂O₄@CMC and investigate its effectiveness in removing CIP during the photocatalytic process. The authors successfully synthesized the nanobiocomposite ZnFe₂O₄@CMC via hydrothermal method. Initially, Fe(NO₃)₃·9H₂O and Zn(NO₃)₂·6H₂O were dissolved in a 2:1 ratio in 100 mL of deionized water. Then 0.5 g of carboxymethylcellulose (CMC) and 6 g of NaOH was being gradually added over an hour to obtain a suspension with pH = 12. After 30 min, a suspension of dark brown color was obtained and placed in the oven at 160 °C for 20 h. The acquired precipitate was washed and dried at 60 °C for 2 h. TEM image of ZnFe₂O₄@CMC as a prepared photocatalyst for degradation of CIP is shown in Figure 9c. The photocatalytic activity of ZnFe₂O₄@CMC was evaluated by studying the effect of reaction time (20–120 min),
the initial concentration of CIP (5–30 mg/L), pH (3–11), the dose of photocatalyst (0.1–0.5 g). The optimal conditions for maximum removal efficiency in synthetic (87%) and natural (79%) samples were: pH = 7, initial CIP concentration—5 mg/L, photocatalyst dose—0.3 g and irradiation time—100 min. Kinetic studies have shown that photocatalytic degradation of CIP is accompanied by pseudo-first-order kinetics. Mechanism for photodegradation of CIP by nano ZnFe$_2$O$_4$@CMC is shown in Figure 9d. The new magnetic nanobiocomposite ZnFe$_2$O$_4$@CMC has demonstrated good chemical stability and reusability after five cycles.

Saied Taghavi Fardood et al. [160] reported non-toxic, cheap, and eco-friendly route using tragacanth gel to synthesize superparamagnetic nanoparticles of magnesium ferrite (MgFe$_2$O$_4$). The MgFe$_2$O$_4$ NPs exhibits excellent photocatalytic properties for Malachite green dye removal (98%) when exposed to visible light. The MgFe$_2$O$_4$ catalyst demonstrate good reusability during six cycles and can be easily removed from solution by magnetic separation.

Gayathri Manju et al. [161] prepared nickel-copper ferrite nanoparticles [Cu$_{1-x}$Ni$_x$Fe$_2$O$_4$ ($x = 0, 0.5, 1, 1.5$)] via combustion method using Aloe barbadensis extract as a “green” reducing agent. XRD patterns confirmed the formation of compositions with a cubic spinel structure and a crystallite size of 52 nm shrank to 29 nm after adding nickel to copper ferrite and to 35.85 nm for nickel ferrite. Measurements of magnetization received at room temperature indicated mild ferromagnetic behavior and saturation magnetization, and the coercivity value increased with nickel substitution. A study of antibacterial activity against Bacillus subtilis, S. aureus, Klebsilla pneumonia and E.coli was performed using diffusion method. The results showed increased activity when adding nickel to copper ferrite.

Atrak et al. [162] synthesized Mg$_{0.5}$Ni$_{0.5}$Al$_x$Fe$_{2-x}$O$_4$ ($x = 0.0, 0.5, 1, 1.5$) spinel ferrites by a “green” sol-gel method using tragacanth gel (Figure 10a,b). It was shown that the crystallite size decreased with increasing concentration of Al$^{3+}$. Studies of the optical bands energy gap in the samples show that the value of the band gap increases from 2.55 to 2.67 eV due to increase in the dosage of Al$^{3+}$. Photocatalytic activity was evaluated during the degradation of the direct blue 129 (DB129) dye as a model reaction of environmental pollution when exposed to visible light. Experimental results confirmed a direct relation between photocatalytic activity and the amount of Al: a catalyst with $x = 1.5$ illustrates better degradation (94%) than a catalyst with $x = 1$ (88%) and $x = 0.5$ (79%).

Mahajan et al. [153] synthesized CoFe$_2$O$_4$ NPs and Ag$_x$Co$_{1-x}$Fe$_2$O$_4$ NPs (where $x = 0, 0.005, 0.01, 0.02$) via sol-gel auto combustion method using “green” and chemical synthesis with extracts from tulsa seed (Ocimum sanctum) and garlic cloves (Allium sativum). The XRD analysis confirmed that the prepared samples were crystalline and had a cubic spinel (inverse) structure. Magnetic measurements at room temperature showed that the saturation magnetization values for samples obtained using tulsa seed extract (49.72 emu/g) and chemical synthesis (49.95 emu/g) were significantly higher than those obtained using garlic extract (28.89 emu/g) (Figure 10c). FE-SEM micrograph of Ag 1% doped chemically synthesized CoFe$_2$O$_4$ nanoparticles is shown in Figure 10d. The samples demonstrated good antibacterial activity: they were more effective against gram-positive than against gram-negative bacteria, mainly due to the difference in the bacterial cell wall.

Madhuikara Naik et al. [152] prepared nanostructured Zn-doped cobalt ferrites Zn$_x$Co$_{1-x}$Fe$_2$O$_4$ ($x = 0.0$ to 0.6) via combustion method using cheese as “green” fuel. X-ray diffraction patterns reveal that these nanomaterials have a crystallite size in the range of ~12–21 nm with an inverse cubic spinel structure. Figure 11a shows CIE diagram of Zn$_x$Co$_{1-x}$Fe$_2$O$_4$ NPs. The vibrational stretching modes of the tetrahedral (582 cm$^{-1}$) and octahedral (385 cm$^{-1}$) sections (metal-oxygen bonds) were confirmed by IR spectra. Photodegradation studies of the obtained nanoparticles were examined during degradation of Congo red and Evans blue dyes under visible light (Figure 11b). Pure CoFe$_2$O$_4$ nanoparticles and Zn-doped CoFe$_2$O$_4$ NPs were examined against both gram-positive (S. aureus) and gram-negative (S. typhi) bacteria. Gram-negative bacteria Salmonella typhi exhibit high antibacterial activity in a zone of inhibition of Zn-doped CoFe$_2$O$_4$ (22 mm), compared
with pure CoFe$_2$O$_4$ (16 mm). The obtained nanoparticles are suitable for optoelectronic, photocatalytic and pharmaceutical applications.

Moradnia et al. [154] obtained MgFeCrO$_4$ magnetic nanoparticles of spinel structure via a “green” sol-gel synthesis method. Tragacanth gel was used to provide a comfortable, natural and cheap sol-gel method that does not contain surfactants and organic solvents. X-ray diffraction patterns confirmed the formation of spinel cubic magnetic MgFeCrO$_4$ nanoparticles. Figure 11c shows magnetic hysteresis loop of spinel MgFeCrO$_4$ nanoparticles, and Figure 11d shows TEM image pattern of MgFeCrO$_4$ MNPs. The unique photocatalytic activity of magnetic MgFeCrO$_4$ nanoparticles for rapid degradation of direct black 122 (DB122) dye in aqueous solution under visible light was studied. The photocatalytic activity of the nanocatalyst (MgFeCrO$_4$) is achieved due to the synergistic effect between Mg, Fe and Cr in the spinel structure. This nanocatalyst is heterogeneous (insoluble in water) and stable during photodegradation. The results showed that 96% of the DB122 dye was degraded in only 60 s. The degradation kinetics of DB122 are consistent with the pseudo-first-order kinetic model. Magnetic MgFeCrO$_4$ nanoparticles show excellent reusability for DB122 dye degradation, as the photocatalyst did not show a significant reduction in its activity even after four applications. The synthesized magnetic nanoparticles have a promising potential for use in electrical and optical systems, cosmetology, ecology, etc.

**Figure 10.** (a) Magnetic hysteresis loops of spinel Mg$_{60.5}$Ni$_{0.5}$Al$_x$Fe$_{2-x}$O$_4$ (a: $x = 0.5$; b: $x = 1$; c: $x = 1.5$) obtained using tragacanth gel (Adapted with permission from [162], Elsevier, 2019); (b) TEM images of Mg$_{60.5}$Ni$_{0.5}$Al$_x$Fe$_{2-x}$O$_4$ MNPs ($x = 0.5$) (Adapted with permission from [162], Elsevier, 2019); (c) magnetic properties and (d) FE-SEM micrograph of Ag 1% doped CoFe$_2$O$_4$ nanoparticles, synthesized using green extract of garlic and tulsi seed (Adapted with permission from [153], Elsevier, 2019).
Figure 11. (a) CIE diagram of ZCF NPs and (b) possible reaction mechanism for the photocatalytic degradation of CR and EB dyes over ZCF NPs (Adapted with permission from [152], Elsevier, 2019); (c) Magnetic hysteresis loop and (d) TEM image of MgFeCrO₄ MNPs (Adapted with permission from [154], Elsevier, 2020).

5. “Green” Synthesis of Metal Nanoparticles

5.1. Silver Nanoparticles

Out of all noble metal nanoparticles, silver nanoparticles (AgNPs) are widely used in various fields, such as optic materials [163], photocatalysts [164–169], biomedicine [170–176], etc. For example, the general use of AgNPs in biomedicine can be attributed to their potent antibacterial properties against a wide range of bacteria. Various methods are used effectively to produce large numbers of AgNPs. However, these methods remain relatively expensive and usually require the involvement of certain harmful substances. Therefore, the development of “green” methods for obtaining silver nanoparticles is very important [177]. Recently, the obtaining of silver NPs using biological processes has attracted much attention. The use of plants and plant extracts is one of the most desirable methods of “green” biological synthesis due to their rich biologically active metabolites [178–186].

Aygün et al. [187] synthesized silver nanoparticles (AgNPs) using reishi mushroom extract (Ganoderma lucidum). 20 mL of mushroom extract was diluted to 100 mL by adding distilled water. Later, 15 mg of AgNO₃ salt was added to the mixture, which was then stirred magnetically until Ag⁺ ions reduced to Ag⁰ ions (transition from clear to brown-red color of the solution). In the UV-visible spectrum, a wide absorption peak between 400–460 nm was detected, indicating the presence of AgNPs. TEM images proved the nanoparticles had spherical shape with a diameter of 15–22 nm (Figure 12a). Antioxidant, antifungal and antimicrobial (against S.aureus, E.coli, P.aeruginosa, L.Pneumophila, C. albicans strains) activity of AgNPs were studied.

De Aragão et al. [188] offered a simple method of “green” route of AgNPs synthesis, using a polysaccharide extracted from red algae Gracilaria birdiae. AgNPs ranged between 20.2 nm and 94.9 nm (Figure 12b). AgNPs were examined for antimicrobial activity against Escherichia coli and Staphylococcus aureus. The obtained results prove that the silver nanoparticles, synthesized along with the polysaccharide, can be used for drug delivery.
Khatami et al. [189] synthesized AgNPs using dried grass. The average size of AgNPs was 15 nm (Figure 12c). It was investigated that such NPs demonstrate the antitumor, antibacterial and antifungal activity. When the concentration of AgNPs increases to five µg/mL, an inhibitory effect on the cancer cells growth is achieved, the survival of cancer cells is reduced by approximately 30%.

Saha et al. [190] presented a “green” eco-friendly method of AgNPs biogenic synthesis using *Gmelina arborea* fruit extract. The mixture of fruit extract and AgNO$_3$ was being heated at 60 °C along with continuous magnetical stirring at 1000 rpm. Within 5 min, the color of the solution changed from clear to yellowish. The formation of AgNPs was investigated via UV-visible spectra over the same period. A blank probe was prepared by taking 30 mL of distilled water instead of silver nitrate. The amount of added fruit extract ranged from 0.1 to 1.0 mL. TEM analysis proved that the AgNPs are stable, with spherical shape, and particle sizes ranging from 8 to 32 nm. The AgNPs have been tested as “green” catalyst in Methylene blue dye degradation and demonstrated excellent catalytic properties.

M.R. Bindhu et al. [182] TEM image of AgNPs synthesized using *Moringa oleifera* flowers (Figure 12d). Transmission electron microscopy analysis indicated the formation of nanoparticles with spherical shape and size of 8 nm. Synthesized “green” AgNPs suppressed the growth of *Klebsiella pneumonia* and *Staphylococcus aureus* and effectively sensed Cu$^{4+}$ ions.

Nouri et al. [191] synthesized ultra-small AgNPs using *Mentha aquatica* leaf extract. Phytochemicals from the extract can reduce Ag$^+$ to Ag$^0$ and form nanoparticles. The synthesis of AgNPs was carried out from AgNO$_3$ solution and at different pH (9, 9.5, 10 and 10.5), volume ratio of AgNO$_3$ solution to the extract (0.1: 0.9, 0.3: 0, 7, 0.5: 0.7: 0.3, 0.9: 0.1 mL/mL), temperature (25, 40, 70, 90 °C) and ultrasound power (50, 100, 150 and 200 W). 0.5 mL of leaf extract was added to 19 mL of water, the pH was adjusted by adding 0.2M K$_2$CO$_3$ solution. Next, 0.5 mL of aqueous AgNO$_3$ solution (100 mM) was added dropwise, while continuously stirring magnetically or subjecting to ultrasound with a high-power ultrasonic generator tool equipped with a titanium tip. The formation of AgNPs through the reduction of Ag$^+$ to Ag$^0$ was monitored over different amounts of time via UV-spectroscopic analysis. To prevent unnecessary photochemical reactions, the vessel was covered with aluminum foil. Furthermore, all glassware was washed with a mixture of solutions of HCl and HNO$_3$ (HCl:HNO$_3$ = 3:1 v/v), and dried at 100 °C. The obtained AgNPs were washed and centrifuged at 16,000 rpm. The effective synthesis parameters were fully optimized to achieve small size of nanoparticles (8 nm) (Figure 12e) with superior antibacterial properties. In particular, the values of the minimum inhibitory concentration for ultrasonically synthesized AgNPs against *P. aeruginosa*, *E. coli*, *B. cereus* and *S. aureus* were 2.2, 58, 20 and 198 µg/mL, respectively. In addition, those AgNPs have shown significant catalytic activity for the removal of various types of dyes, which are environmental contaminants.

Ravichandran et al. [167] performed a “green” synthesis of AgNPs via bioreduction of silver nitrate using an *Parkia speciosa* leaf extract. The change of the solution color to brown indicated the biological reduction of AgNO$_3$: 1 mL of 0.01M AgNO$_3$ solution was added to 1 mL of *Parkia speciosa* leaf extract in a 10 mL volumetric flask, which was filled to 10 mL with deionized water and kept at room temperature for 24 h. The synthesized AgNPs were centrifugated at 10,000 rpm. The synthesized AgNPs demonstrated maximum absorption at 410.5 nm. SEM and TEM (Figure 12f) analyses revealed the average size of AgNPs (31 nm and 35 nm, respectively). The synthesized AgNPs demonstrate good photocatalytic (in Methylene blue dye degradation), antimicrobial (against *Bacillus subtilis*, *Pseudomonas aeruginosa*, *Staphylococcus aureus* and *Escherichia coli*) and antioxidant activity. Synthesized AgNPs can be used in various fields, such as water purification, biomedicine, biosensors and nanotechnology.
ids, flavonoids, saponins, anthraquinones, and steroids. The plasmon resonance peak was at 350 nm, and such functional groups as hydroxyl, carboxyl, alkyl halide, phenolic, amine, carbonyl, and amide were important for bioreducing and closing silver ions into nanoparticles. The analysis showed that silver is the main element, and nanoparticles have an irregular shape and size of 12 nm. The obtained creams were stable, cosmetically attractive with satisfactory pH, viscosity and application. Silver nanoparticles based on aqueous extracts of Nauclea latifolia bioreduce silver nitrate to AgNPs. Phytochemical evaluation of crude Nauclea latifolia extracts showed the presence of alkaloids, terpenoids, flavonoids, saponins, anthraquinones, steroids, and tannins. The plasmon resonance peak was at 350 nm, and such functional groups as hydroxyl, carboxyl, alkyl halide, phenolic, amine, carbonyl, and amide were important for bioreducing and closing silver ions into nanoparticles. The analysis showed that silver is the main element, and nanoparticles have an irregular shape and size of 12 nm. The obtained creams were stable, cosmetically attractive with satisfactory pH, viscosity and application. Silver nanoparticles based on aqueous extracts of Nauclea latifolia and its cream composition showed strong antimicrobial and antioxidant activity.

5.2. Gold Nanoparticles

Gold nanoparticles (AuNPs) are considered the most attractive among noble metal nanoparticles, because of their potential use in catalysis, optics, biomedicine, etc. The widespread application of AuNPs has aroused considerable interest for new AuNPs synthesis methods, avoiding hazardous chemicals [53,177,193–196]. Alternative methods, including “green” synthetic approaches to produce AuNPs, are important for maintaining sustainable development.

Odeniyi et al. [192] investigated the phytochemical, antioxidant and antimicrobial potential of aqueous and methanolic extracts of Nauclea latifolia fruit and their application in the biosynthesis of AgNPs and for cold cream formulations. The extracts were used to bioreduce silver nitrate to AgNPs. Phytochemical evaluation of crude Nauclea latifolia extracts showed the presence of alkaloids, terpenoids, flavonoids, saponins, anthraquinones, steroids, and tannins. The plasmon resonance peak was at 350 nm, and such functional groups as hydroxyl, carboxyl, alkyl halide, phenolic, amine, carbonyl, and amide were important for bioreducing and closing silver ions into nanoparticles. The analysis showed that silver is the main element, and nanoparticles have an irregular shape and size of 12 nm. The obtained creams were stable, cosmetically attractive with satisfactory pH, viscosity and application. Silver nanoparticles based on aqueous extracts of Nauclea latifolia and its cream composition showed strong antimicrobial and antioxidant activity.
discarded and oven dried at 60 °C. The Fe₃O₄/Au NPs demonstrate magnetic properties (Figure 13a) and average diameter is around six nm (Figure 13b). The anticancer activity of Fe₃O₄/Au NPs may be promised candidates for cancer treatment and other biomedical applications.

![Figure 13.](image_url)

Figure 13. (a) Vibrating sample magnetometer plots of Fe₃O₄ and Fe₃O₄ coated with Au nanoparticles (Adapted with permission from [194], Elsevier, 2019); (b) HR-TEM image of Fe₃O₄/Au nanoparticles (Adapted with permission from [194], Elsevier, 2019); (c) UV-Visible spectrum of Cr@AuNPs. The inserted figure shows the color changes of Cr@AuNPs, (i) extract, (ii) chloroauric acid and, (iii) synthesized Cr@AuNPs (Adapted with permission from [198], Elsevier, 2019); (d) XRD pattern of Cr@AuNPs (Adapted with permission from [198], Elsevier, 2019).

Manikandakrishnan et al. [198] reported using Caulerpa racemosa (Cr) to synthesize gold nanoparticles (AuNPs). C. racemosa extract mixed with HAuCl₄ solution under stirring (24 h) and the Cr@AuNPs were obtained (Figure 13c). The presence of phytochemical components in the aqueous extract of C. racemosa was confirmed by IR spectroscopic analysis. The size of Cr@AuNPs ranged from 13.7 to 85.4 nm. XRD pattern of Cr@AuNPs is shown in Figure 13d. The synthesized Cr@AuNPs effectively controlled the growth of human colon adenocarcinoma cells (HT-29), and demonstrated IC₅₀ of 20.84 µg/mL. “Green” synthesized Cr@AuNPs had a non-toxic effect on Artemia nauplii, even at high concentrations (100 µg/mL).

Shabestarian et al. [199] developed a “green”, eco-friendly, fast and simple synthesis of AuNPs using aqueous Sumac extract. The solution of HAuCl₄ mixed with aqueous Sumac extract and the purple solid was obtained. The TEM analysis revealed that the bio-formed AuNPs have a spherical morphology with an average size of 20 nm. The antioxidant activity of the bio-formed AuNPs has been tested. It was concluded that such AuNPs are promising for biomedical applications.
5.3. Platinum Nanoparticles

Platinum nanoparticles (PtNPs) have been widely used in oxidation and hydrogenation reactions in petrochemical industry, due to their large surface and many other characteristics that encourage to synthesize PtNPs for catalytical applications. Therefore, there is a high demand for synthesis of PtNPs using of eco-friendly materials [177,200–205].

Kumar et al. [201] used Xanthium strumarium leaf extract for the biosynthesis of PtNPs. TEM analysis confirmed the formation of PtNPs with an average size of 22 nm, and SEM analysis revealed that those PtNPs have a cubic and rectangular shape and smooth surface. The nanoparticles demonstrated huge cytotoxic effect on HeLa cell lines, thus can be used for biomedical applications.

Dobrucka [203] presented „green” synthesis of PtNPs using Fumariae herba extract. TEM analysis showed hexagonal and pentagonal shapes of synthesized PtNPs with a diameter of 30 nm. PtNPs demonstrated high catalytic activity during reduction of violet crystal and methyl blue dyes.

Al-Radadi et al. [200] performed a “green” synthesis of PtNPs using an extract solution of dates (biodegradable surfactants) in order to learn about their effect on various cancer cells. Aqueous extracts of solutions from Ajwa and Barni dates behave as stabilizers and reductants in the synthesis of PtNPs in natural environment conditions. The size of obtained PtNPs was small, in the range of 1.3-2.6 nm. Furthermore, the change of pH in the reaction affected the size of nanoparticles. Antitumor activity of PtNPs was tested for various cancer cells: carcinoma cells (HCT-116), breast cancer cells (MCF-7) and hepatocellular carcinoma (HePG-2). Antibacterial activity of PtNPs against Escherichia coli and Bacillus subtilis bacteria was study and were found that PtNPs are able to inhibit the growth of E. coli and B. subtilis.

5.4. Palladium Nanoparticles

Palladium nanoparticles (PdNPs) are being used in various unique applications, including sensors and active membrane catalysts, thanks to their catalytic properties and hydrogen affinity. PdNPs are synthesized using a variety of wet chemical approach, such as chemical reduction, electrochemical and polyol methods [177].

„Green” synthesis of PdNPs for catalysis and biological applying is of enormous interest [206–213]. Lentinan (LNT) can be a fine stabilizing and reducing agent for replacing complex plant extracts. Han et al. [214] showed a simple “green” method of PdNPs synthesis using LNT as a stabilizer to achieve a smaller Pd\textsubscript{n}-LNT nanoparticles (2.35–3.32 nm), as well as a higher catalytic activity for the 4-nitrophenol reduction, comparing to other known catalysts. Furthermore, Pd\textsubscript{n}-LNT NPs demonstrated insignificant cell cytotoxicity along with good antioxidant activity.

Celebioglu et al. [215] synthesized PdNPs loaded on cyclodextrin (CD) nanofibers. Cyclodextrin acted as a reductant and catalyzed the PdNPs formation upon reduction from Pd\textsuperscript{2+} to metallic Pd\textsuperscript{0}. The results of the study confirmed the presence of homogeneously distributed polycrystalline PdNPs (3–5 nm) in the entire nanofiber matrix and shown the existing of a larger fraction of the metal Pd\textsuperscript{0} atom due to effective reduction of Pd\textsuperscript{2+} by CD molecules. Catalytic properties of PdNPs were evaluated through reduction of n-nitrophenol to n-aminophenol, resulting in high catalytic activity of nanofibers.

Amrutham et al. [216] synthesized nano-sized PdNPs via a new, single-stage and cheap method with high yield—microwave irradiation using water-soluble Neem wood resin as a stabilizer and reductant. The resin was purified and washed thoroughly with double distilled water. An aliquot of 50 mL of an aqueous H\textsubscript{2}PdCl\textsubscript{4} solution (5 × 10\textsuperscript{-4} M) was added to 50 mL of a 1% resin aqueous solution. The reaction took place in a microwave oven at 320 W for 10 min. The average particle size was about four nm. Catalytic activity of PdNPs was evaluated spectrophotometrically through reduction of Rhodamine 6 G dye using NaBH\textsubscript{4}. PdNPs, stabilized with Neem wood resin, demonstrated excellent catalytic activity for Rhodamine 6 G reduction (18 min). The reaction showed pseudo-first order kinetics, and the obtained rate constant equaled 0.1875 min\textsuperscript{-1}. 
6. Future Perspectives

Thus, green chemistry is an ecological branch of chemistry that aims to minimize the use of toxic and hazardous chemicals. There are several main ways to maximize the implementation of the principles of “green” chemistry in all spheres of life:

(1) to create various chemical associations, organizations and institutes, whose missions will be studying of cleaner reactions, products and processes;
(2) to promote “green” chemistry among universities and research laboratories in order to develop economically sustainable technologies for clean production;
(3) to introduce methods of “green” synthesis of chemicals in industrial enterprises;
(4) to train future scientists in universities, who in the future will solve regional and global environmental problems (nowadays, most industrial developments are related mainly to economic efficiency, rather than environmental friendliness of processes);
(5) to ensure environmental protection at the legislative level;
(6) to use innovative alternative methods of minimal production of undesired chemicals in order to preserve human health and reduce harmful effects on the environment, namely: use alternative (renewable) sources of raw materials; use less hazardous reagents; use alternative solvents (ionic liquids, water, etc.) during the synthesis of organic matter; use “green” catalysts that affect energy consumption and reduce the production of unwanted by-products and waste; minimize energy consumption at every stage of the industrial process.

In future, “green” chemistry should be applied to almost every sector of the business—food industry, energy, plastics, medicines, cosmetics, cleaning products, etc., and therefore it will play an important role in the industry development.

7. Conclusions

This review reveals basic principles of “green” chemistry and “green” methods for obtaining metal/metal oxide nanoparticles using bioresources. The most common method for obtaining these nanoparticles is a “bottom-up” approach, using various organic solvents, toxic chemicals and non-ecological reagents under high pressure and temperature. Therefore, alternative, cheap and safe methods are necessary. “Green” synthesis prevents pollution during initial stages of chemical processes and reduces the negative impact on the environment and human health. Many interesting biological methods using plants, algae, fungi, yeast, bacteria, viruses, etc., have been developed. Among various “green” resources, plants are the best source of precursors for the synthesis of metal/metal oxide nanoparticles, due to their simplicity, non-toxicity and availability. Such parameters as total antioxidant capacity and total protein content affect the suitability of plants. It is important to study physical and chemical properties, stability and activity of “green” nanoparticles in order to further improve their practical application. This review involves using sol-gel method, hyperthermal method, auto-combustion, etc., for the synthesis of magnetic nanoparticles. Various extracts of plants, including leaves of Lagenaria siceraria, Aloe vera, Averrhoa carambola, husk of Punica Granatum, roots of Mimosa pudica, Chromolaena odorata, etc., are frequently used as reductants. The separate section is dedicated to “green” synthesis and the use of spinel magnetic nanoparticles (in particular, superparamagnetic \( \text{Fe}_3\text{O}_4 \) NPs), which attracted the attention of researchers due to their unique properties: bio-degradability, biocompatibility and non-toxic effect. The biocompatibility and high saturation magnetization of naturally stabilized magnetic nanoparticles allows for their various biological applications. Methods of “green” synthesis of silver, gold, platinum and palladium nanoparticles are also described. For reduction and stabilization during the synthesis of AgNPs, extracts of reishi mushroom (Ganoderma lucidum), red algae (Gracilaria birdiae), Gmelina arborea fruit, Nauclea latifolia fruit, Mentha aquatica leaves and Parkia speciosa leaves have been used. For the synthesis of AuNPs, Juglans regia green husk extract, Caulerpa racemose extract and Sumac aqueous extract have been used. For the synthesis of PtNPs, leaf extracts of Xanthium strumarium and Fumariae herba have been used. For the synthesis of PdNPs, Neem wood resin extract has been used. The obtained nanoparticles are
suitable for optoelectronic, photocatalytic and pharmaceutical applications. The synthesis of metal/metal oxide nanoparticles via “green” approach allows for obtaining nanoparticles with specified sizes and improved morphology. These new “green” technologies can radically reduce environmental pollution and risk to human health.

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