



Contributions of Nano-Nitrogen Fertilizers to Sustainable Development Goals: A Comprehensive Review

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Abstract: Nano-nitrogen fertilizers (NNFs) have emerged as a promising technology in the field of agriculture, offering potential solutions to improve nutrient uptake efficiency, enhance crop productivity, and reduce environmental impacts. NNFs showed superior characteristics and performance on crops and, therefore, became a potential alternative to conventional nitrogen (N) fertilizers. These fertilizers enhance plant uptake while simultaneously reducing environmental losses. For example, a hydroxy appetite-based urea NNF extended the N release for 112 days, which could cover the N demand of many perennial crops, thus reducing losses. The reported NNFs in this review increased the yield by 10–80% compared to conventional fertilizers. Additionally, their small particle size increases crop acclimation and decreases the application rate. With all these beneficial traits of NNFs, they potentially contribute to achieving Sustainable Development Goals (SDGs). This review article summarizes the materials used in NNF formulation, methods of preparing NNFs, and their crop responses. Also, it highlights the limitations identified in the research studies and provides research recommendations for the future. Further, it provides a critical assessment of the current state of NNFs and their prospects for revolutionizing modern agriculture to attain SDGs.

Keywords: crop responses; environmental impacts; nanoparticles; smart fertilizers; synergist



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1. Introduction

The ever-growing population has a substantial impact on food demand and necessitates higher agricultural production [1]. With limited arable land, increasing the agricultural input rates, including nitrogen (N) fertilizers, to obtain higher productivity is seen as a successful way to address the growing food demand [2]. Nitrogen is a major nutrient for plants that primarily influences vegetative growth and crop yield [3]. Although higher levels of N fertilizer are applied with the goal of achieving higher yields, only a small fraction is actually taken up by the plants, while a significant portion is lost through processes such as leaching, runoff, and gaseous emissions, namely N_2O , NO, and NH_3 [4,5]. With the main focus on increasing productivity, the negative environmental consequences of increasing the application rate or excessively applying N fertilizers have been often overlooked in the past [6]. However, after several studies showed the detrimental effects of higher N application rates on soil health, water quality, and ecosystem sustainability, a paradigm shift has taken place toward the sustainable use of N fertilizers in agricultural systems.

The current and future agriculture practices should align with the Sustainable Development Goals (SDGs) established by the United Nations to effectively tackle global challenges such as poverty, hunger, climate change, and environmental degradation. Sustainable agriculture plays a crucial role in achieving the SDGs. Achieving SDGs can be facilitated through the precise application of input resources, particularly N fertilizers. Toward this, several approaches are being used, namely split applications, integrated nutrient management, application closer to the root zone, foliar application, the application of fertilizer after substantial development of crops, and the use of smart fertilizers [6,7]. Of these, smart fertilizers are the latest technology used to minimize nitrogen losses to the environment and increase nitrogen utilization efficiency (NUE).

Smart fertilizers, also known as slow- (SRFs) or controlled-release fertilizers (CRFs), have the ability to release nutrients in a slower manner than conventional fertilizer, and their release pattern matches the crop nutrient demand [8,9]. Several materials have been used in the formulation of SRFs, and nanomaterials have emerged as a recent addition to this repertoire. These materials are within the range of 1–100 nm. Nanomaterials offer unique properties to improve nutrient releases, such as high surface area, high reactivity, and high porosity, thus advancing the field of fertilizer technology [10,11]. Owing to the high surface area to volume ratio, nanomaterials adsorb nutrients higher than bulk materials. Hence, the loading capacity of nanomaterials is very high [12]. Nanofertlizers showed higher absorbance than conventional fertilizers when they were used as foliar applications [13]. Furthermore, nanoparticles are applied as synergists along with fertilizers to increase crop performance [10,14]. Given the benefits of nanofertilizers, they have the potential to be a significant player in achieving SDGs as they can minimize the application rate, frequency of application, losses to the environment, and control greenhouse gas emissions. Although several studies reported the significance of nanofertlizers in agriculture, a dedicated study for nano-nitrogen fertilizers is unavailable. Therefore, this study summarizes the materials used for formulating NNFs, methods employed in preparing NNFs, and their responses to crops.

2. Materials Used in Nanoparticle Preparation

2.1. Clay Minerals

Clay minerals are used in nanofertilizers due to their unique properties such as high cation exchange capacity (CEC), porous structure, high surface-to-volume ratio, colloidal property, and ease of modification. Additionally, they are readily available and cheap materials compared to their counterparts. Bentonite [15], attapulgite [16], kaoline [17], and glauconite [18] are a few of the clay minerals used for the formulation of NNFs.

Bentonite is an aluminum phyllosilicate clay that is primarily composed of montmorillonite, a member of the smectite group of minerals [19]. It is derived from volcanic ash deposits that have undergone weathering and transformation over time. The lamellar structure of bentonite consists of two silica tetrahedral sheets with an alumina octahedral sheet sandwiched in between. The sheet structure of attapulgite contains interlayer spaces that are occupied by various cations. These cations have the ability to be replaced by other cations [20]. Therefore, this provides the opportunity to incorporate several macro and micronutrients with this clay. The CEC of this clay ranged between 60 and 150 meq 100 g⁻¹ of soil [21].

Several NNFs were developed using bentonite as a raw material [11,15]. In a study, Umar et al. [11,15] developed an SRF by coating urea with Zn-fortified nano-bentonite (Table 1). To enhance the presence of Zn^{2+} on the active sites of bentonite, nano-bentonite was treated with varying concentrations of a ZnSO₄ solution. Subsequently, urea was coated using two different methods: firstly, by employing vegetable oil and nano-bentonite (referred to as ZnBenVegU), and secondly, using stearic acid, paraffin oil, Ca(OH)₂, and nano-bentonite (referred to as ZnBenParU). The results of a soil incubation study (Table 1) indicated that the release of urea was effectively controlled for up to 10 days with ZnBenVegU and up to 15 days with ZnBenParU. The authors propose that the network structure of bentonite potentially increases the distance water must traverse, thereby reducing urea dissolution. Liu et al. [11,15] developed an SRF by reacting biochar together with bentonite and polyvinyl alcohol (PVA) and impregnated the urea into it (Table 1). Through a reaction between the -OH group of bentonites and biochar along with polyvinyl alcohol (PVA), new bonds were formed.

Fertilizer	N Source	Nutrients	NP Used	NP Preparation or Modification Method	Binder/Other Components	Method of SRF Preparation	N Release	References
ZnBenVegU	Urea	N and Zn	Zn fortified nano-bentonite	Soil-gel	Vegetable oil	Coating	10 days	[15]
ZnBenParU	Urea	N and Zn	Zn fortified nano-bentonite	Soil-gel	Stearic acid, paraffin oil, and paraffin wax	Coating	15 days	[15]
QAL-Ben-U	Urea	Ν	Bentonite	Soil–gel	Quaternary ammonium lignin (QAL)	Matrix and coating	N/A	[22]
Nano-biochar SRF	Sodium nitrate	N, P, K, Ca, and micronutrients	Nano-biochar	Physical crushing	N/A	Impregnation	>10 days	[23]
U-CAM	Urea	N, Fe, and Ca	Carboxylated nanocellulose (CNF)	Catalytic oxidation	Hydrogel	Matrix	>30 days	[24]
BNC fertilizer	Sodium nitrate	N, Ca, P, K, Mg, and micronutrients	Nano-biochar	Physical crushing	N/A	Impregnation	>14 days	[25]
WNLCU	Urea	Ν	Attapulgite (HA)	High-energy electron beam (HEEB) irradiation	Sodium polyacrylate (P) and polyacrylamide (M)	Matrix	66% lower than control	[16]
WNLCN	Ammonium chloride	Ν	Attapulgite (HA)	High-energy electron beam (HEEB) irradiation	Sodium polyacrylate (P) and polyacrylamide (M)	Matrix	90% lower than control	[16]
Loss control urea (LCU)	Urea	Ν	Attapulgite	Irradiated by high-energy electron beam and O ₃ treatment	Polyacrylamide (P)	Matrix	50% lower than urea	[26]
Coated urea	Urea	Ν	Kaoline and Polystyrene-starch	Ultra-high- speed cutting and semi-emulsification	None	Coating	N/A	[17]
Kao-urea	Urea	Ν	Kaolin	None	Chitosan	Matrix	>30 days	[27]
Kao-urea	Urea	Ν	Kaolin	Milling	None	Matrix	>7 days	[28]
Gal-ADP	Ammonium dihydrogen phosphate (ADP)	N, K, P, and other micronutrients	Glauconite	Chemical and mechanochemical method	Na ₂ CO ₃ as an extender	Matrix	>56 days	[18]
HA-POL-urea	Urea	N, K, and P	Hydroxyapatite (HDA)	Sol—gel	Cellulose fiber and polyacrylamide	Matrix	112 days	[29]

Table 1. The nanoparticle-coated slow-release N fertilizers (NSRNFs) and their method of development and response on crops.

Fertilizer	N Source	Nutrients	NP Used	NP Preparation or Modification Method	Binder/Other Components	Method of SRF Preparation	N Release	References
Zeo-AN	Ammonium nitrate (AN)	Ν	Zeolite (surface-modified)	Hydrothermally synthesized	None	Surface carrier	35% lower than CF	[30]
Nano Zn-MAP and nano-Zn-urea	Monoammonium phosphate (MAP) and urea	Ν	ZnO	N/A	Water	Coating	N/A	[31]
Zeolite	Sodium nitrate	N and other macro- and micronutrients	Zeolite	Co-precipitation method	None	Surface carrier	>7 days—water and >16 days—soil	[32]
Zn-MAP and Zn-urea	Monoammonium phosphate (MAP) and urea	N and Zn	ZnO	None	Water	Coating	N/A	[33]

The findings of this study demonstrated that bentonite and urea were successfully incorporated within the cavities and channels of biochar and underwent polymerization with PVA. This impregnation and polymerization process effectively slowed down the dissolution of urea in water for a period of up to 42 days. In another study, urea was intercalated with quaternary ammonium lignin (QAL) modified nano-bentonite and then mixed with sodium alginate to form NNFs [22] (Table 1).

Attapulgite, also known as palygorskite, is a hydrated aluminum–magnesium silicate $(Mg,Al)_4(Si)_8(O,OH,H_2O)_{26} \cdot nH_2O)$. This is a naturally occurring nano-clay with a nanorod structure, and the average length and width of this rod are 800–1000 and 30–40 nm, respectively [16]. Attapulgite exhibits a ribbon-layer structure with a 2:1 arrangement in which ribbons are interconnected through the inversion of SiO₄ tetrahedra (Figure 1). This arrangement allows for the formation of channels and tunnels [34], which could accommodate foreign materials or other minerals. The cation exchange capacity of attapulgite ranged between 11 and 33 meq 100 g⁻¹ of soil [35].



Figure 1. A schematic diagram of nanosheet preparation from natural attapulgite [34].

Various NNFs were created by combining modified attapulgite with urea (WNLCU) and ammonium chloride (WNLCN), along with the addition of sodium polyacrylate (P) and polyacrylamide (M) [16] (Table 1). Sodium polyacrylate has high water retention ability and polyacrylamide forms a stronger bond with the nanorod structure of attapulgite. Therefore, the addition of these compounds enhances the network structure of the matrix and extends the nutrient release by 66 and 90% more than the control treatment (Table 1). In a study, modified attapulgite was mixed with polyacrylamide (P) and then loaded into urea or ammonium chloride (NH₄Cl) to obtain loss control urea (LCU) or loss control NH₄Cl (LCN) [26]. This forms a 3D skeleton structure that is mainly formed by modified attapulgite, and P and urea also take part in the structure (Figure 2). The morphology of the 3D skeleton Is pH dependent, and more H-bonds form when pH increases. These fertilizers were reported to decrease N leaching losses by 50% compared to urea and NH₄Cl (Table 1).

Kaolin, which is also referred to as china clay, is a clay mineral primarily composed of kaolinite. It possesses a white color, soft consistency, and fine-grained texture with a smooth feel. Kaolin originates from the weathering and erosion processes of rocks, specifically those rich in feldspar, like granite. The chemical formula of kaolin is $Al_2O_3 \cdot 2SiO_2 \cdot 2H_2O$. The structural arrangement of kaolinite is with one tetrahedral sheet of silica (SiO₄) linked through oxygen atoms to one octahedral sheet of alumina (AlO₆) octahedra [36]. It can interact with other molecules in the inter-layer space.



Figure 2. Scanning electron microscopy (SEM) images of (**a**) a micro/nano network surface of LCU, and (**b**) a transmission electron microscope (TEM) of LCU with a nanorod structure in the skeleton [26].

Kaolin and abandoned plastic (polystyrene) nanocomposites were used as coating materials for SRFs [17] (Table 1). The combination of sodium polyacrylate and polyacrylamide resulted in the formation of a honeycomb structure, as confirmed by SEM images. This structural arrangement led to an increased surface area, enabling effective adsorption and absorption of nutrients, while also functioning as a slow-release fertilizer (SRF). Another NNF was developed by intercalating urea with kaolin and then granulating it with chitosan, as described by Roshanravan et al. [28]. Notably, the native form of kaolin was utilized instead of a modified form in this preparation. As a result, the release of urea was extended beyond 30 days, and the NNF exhibited a controlled release of urea that was 60–77% lower compared to urea (Table 1). Additionally, the study demonstrated that this NNF effectively reduced NH₃ release by 6% compared to urea. Similar urea intercalated NNF with kaolin was produced by co-grinding these raw materials [28]. This study found that increasing milling speed and milling time increased the incorporation of urea into the amorphous structure of kaolin. Further, increasing milling speed and time prolonged the urea release time from the NNF and the best NNF controlled the release up to 7 days (Table 1).

2.2. Minerals

Glauconite is a green-colored mineral belonging to the mica group. It is a hydrous potassium, iron, and aluminum silicate mineral $((K,Na)(Fe,Al,Mg)_2(Si,Al)_4O_{10}(OH)_2)$. It is commonly found in sedimentary rocks, such as sandstones and shales. The structure of glauconite consists of a 2:1 layered arrangement, with an octahedral layer sandwiched between two tetrahedral layers [37]. In preparing NNF, Rudmin et al. [28] found that glauconite was activated using chemical methods, mechanical methods, or a combination of both (Table 1). Ammonium dihydrogen phosphate (ADP) was then loaded into the activated glauconite. Various analyses confirmed that ADP molecules were adsorbed by the surface and meso and macro pores, as well as intercalated between layers of glauconite. In a soil leaching column test, this formulation demonstrated an extended release of nutrients for more than 56 days.

Zeolites are characterized as three-dimensional, microporous, crystalline solids that possess distinct structures composed of aluminum, silicon, and oxygen within their framework. Additionally, the pores of zeolites contain cations and water molecules [32]. Zeolite's high pore density and anion exchange capacity allow for the incorporation of a significant number of anions within its structure. Two different zeolites, synthesized zeolite clinoptilonite (SZC) and synthesized zeolite montmorillonite (SZM), were prepared using silica and aluminum nitrate and loaded with ammonium nitrate (AN) [30]. The nitrogen release of this NNF was 35% lower than AN. Lateef et al. [28] prepared another nano-zeolite

with sodium silicate and ethylene glycol, and then sodium nitrate (SN), urea, and other macro and micronutrients were doped into it. The water incubation and soil leaching study exhibited that these fertilizers could extend the N release for more than 7 and 16 days, respectively (Table 1).

Nano ZnO is extensively utilized as a nanomaterial in various applications, including the formulation of NNFs. Due to its commercial availability, nano-ZnO is easily accessible and can be conveniently used in various applications within the agricultural sector. Milani et al. [31] developed nao-ZnO-coated and regular-ZnO-coated urea/mono ammonium phosphate (MAP) NNFs and compared their characteristics (Table 1). Results revealed that coated urea slightly dissolved and dispersed in the soil compared to the coated MAP, possibly due to the high ionic strength of the urea solution and high pH [38]. This study provides evidence that the solubility of nano-ZnO-coated fertilizers is influenced by the acidity generated by the main nutrient used in the fertilizer. Further, this study revealed that ZnO in coated MAP underwent different speciation like ZnSO₄, Zn(NH₄)PO₄, $(CaZn_2(PO_4)_2 \cdot 2H_2O)$, and $Zn(OH)_2$. Interestingly, the speciation of ZnO differs between nano- and bulk-coated ZnO, suggesting that the speciation of ZnO is dependent on particle size. Indeed, the interactions among ZnO, fertilizers, and soil have a significant impact on the formation of various soluble constituents, which, in turn, affect the controlled release ability of the coatings. These reactions play a crucial role in determining the release kinetics and availability of nutrients from the coated fertilizers in the soil environment. In a previous study by these researchers, it was found that Zn solubility was not significantly influenced by the size of the ZnO particle used for coating urea or MAP [33] (Table 1).

2.3. Nano-Biochar

Nano-biochar has gained significant attention in the field of agriculture due to its diverse applications. It has been recognized for its potential in various areas such as wastewater treatments, soil amendment, environmental remediation, pesticide formulation, and nutrient delivery [39]. Khan et al. [23] produced biochar from wheat straw and prepared the nano-biochar by mechanical grinding (Table 1). An NNF was prepared by impregnating sodium nitrate and other macro- and micronutrients into nano-biochar. The XRD and FTIR analysis confirmed that nutrients impregnated well into the nano-biochar. It controlled the nitrate release for more than 10 days. In a similar method, corn-based nano-biochar was prepared, and nutrients (N, Ca, P, K, Mg) were impregnated [25]. The nutrient release from this NNF extended for more than 14 days (Table 1). In this study, it was observed that nutrient release initially occurred rapidly and then gradually slowed down. This phenomenon can be attributed to the quick release of surface-bound nutrients followed by the slower release of nutrients impregnated into the macro- and mesopores of the material [8].

2.4. Other Nano-Materials

In addition to the materials discussed in this section, other substances, such as nanocellulose [24], and hydroxyapatite [29], were also used for preparing NNFs.

Nanocellulose, derived from cellulose fibers, offers unique properties, including high surface area, biodegradability, and stability, which make it a promising material for NNF formulations [40]. Nanocellulose was prepared from eucalyptus pulp and it was mixed with sodium alginate, FeCl₃.6H₂O (Ferric Chloride; FC), and urea to form a hydrogel as a pH-sensitive NNF [24]. Under microscopic examination, the gel without FC (only nanocellulose) displayed a smooth structure, whereas the introduction of FC resulted in an increase in surface coarseness and roughness. The optimum level of FC is important for the correct level of cross-linking. Increasing the FC content above optimum level led to a weakening of the bonding within the gel matrix. This was evident in the nutrient release in water and soil as well. The NNF with 5%, 10%, and 20% FC extended the 80% of urea release by 3.5, 25, and 5 h, respectively (Table 1). Therefore, this study concluded that 10% of FC and pH 11 was conducive for longer urea retention.

Hydroxyapatite (HA) is a naturally occurring mineral form of calcium apatite, and its chemical formula is $Ca_{10}(PO_4)_6(OH)_2$. Hydroxyapatite finds extensive use in the medical field, specifically in diverse dental applications such as toothpaste, dental fillings, and coatings for dental implants. However, there has been a recent increase in its application within the agricultural sector, especially in developing NNFs [29]. The first nano-HA urea was developed with a ratio of urea:HA of 1:1 using the oven drying method [41] (Table 1).

Another NNF was prepared by incorporating urea into nano-HA at a 6:1 ratio using flash drying [42]. Due to the rapid drying method employed, it was possible to incorporate six parts of urea with one part of nano-HA before phase separation occurred. The researchers noted that surpassing this urea level may result in phase separation during the drying process. This study confirmed the formation of new bonds between urea and nano-HA; however, these bonds were relatively weak, allowing for the release of urea from the matrix to behave as an SRF. Under accelerated leaching conditions, the release of urea was extended up to 3820 s, representing an 86% release, which was 12 times lower than pure urea. Hydroxyapatite, being a slow-release source of phosphorus, is commonly utilized in perennial crops. In the formation of nano-HA, the solubility of P may increase, but it was not investigated in this study (Table 1).

3. Methods of Nanoparticle Formulations and Modifications for Preparing NNFs

There are several nanoparticle formulation and modification methods used in the preparation of NNFs, which are summarized in Figure 3.



Figure 3. Nanoparticle formulation and modification methods.

3.1. Nanoparticle Formulation Methods

There are several methods available for formulating nanoparticles. They are mainly categorized into two methods: top-down and bottom-up approaches. The top-down approach involves breaking down larger particles or bulk materials physically or mechanically to obtain nanoparticles. This approach involves breaking down the starting material into smaller particles through processes such as milling, grinding, or lithography. In the bottom-up approach, nanoparticles are formulated from smaller building blocks or molecular precursors. This approach involves the controlled growth or self-assembly of these building blocks to form nanoparticles with desired properties. The sol–gel method, self-assembly, template-assisted synthesis, microfluidic synthesis, and biomimetic synthesis are a few examples of bottom-up approaches.

3.1.1. Sol-Gel Method

The sol-gel method is widely used in preparing nanoparticles and belongs to bottomup approaches. In this technique, a precursor solution is prepared by dissolving metal hydroxide or metal salts in a solvent (Figure 4). This solution functions as the initiating material for nanoparticle synthesis. Water or a hydrolyzing agent is added to the precursor solution, which generates hydroxyl groups via a hydrolysis reaction. Then, condensation takes place where the hydroxyl groups combine to form new chemical bonds. This forms nanoparticle nuclei within the solution. An aging period will be given to allow the growth of nanoparticles. Finally, nanoparticle suspension dries to remove the solvent entrapped within the pores of the nanoparticle. Based on the drying method, the final product can take two forms: a highly dense xerogel and a more porous aerogel [43].



Figure 4. The generalized sol-gel preparation method.

Several NNFs were synthesized using the sol–gel method. For example, ZnO nanoparticles were prepared by adding and mixing ZnSO₄·H₂O dropwise into a KOH solution [15]. Subsequently, ZnO-NPs were subjected to washing, and ethanol was dried by oven drying at 80 °C. Afterwards, dried ZnO-NPs were calcined at 250 °C in the furnace and ground into powder (Table 1). In preparation for matrix and coated-controlled release urea formulation, the sol–gel method was used [22]. First, quaternary ammonium lignin (QAL) was prepared by mixing epoxypropyl trimethylammonium chloride (ETAC) with lignin dissolved in NaOH. Subsequently, a bentonite clay suspension was mixed with a QAL suspension, and the mixture (QAL-modified bentonite) was freeze-dried. Finally, urea was coated by this mixture in the presence of sodium alginate and a Ca(OH)₂ suspension. In another study, nano-hydroxyapatite (HA) was prepared by adding H₃PO₄ into a Ca(OH)₂ and TX-100 suspension [29]. This mixture was oven-dried at 105 °C and ground into powder. Followed by this, nano-HA was mixed with cellulose fiber and urea, (NH)₄SO₄ and K₂SO₄, to formulate NNFs (Table 1).

3.1.2. Mechanical Attrition

Mechanical attrition is an easy way of synthesizing nanoparticles. This method involves the grinding or milling of bulk materials to reduce the particle size to nanoscale. In mechanical attrition, a high-energy mechanical device, such as a ball mill, rod mill, or ring mill, is used. The repeated collision and impact on the material by the device lead to the fragmentation of the bulk material, ultimately producing nanoparticles [43]. Several studies employed mechanical attrition for preparing the NNFs. In preparation for a nanobiochar SRF, nano-biochar was prepared by grinding the wheat biochar [23]. Similarly, Lateef et al. [25] used a homogenizer to prepare nano-wheat biochar (Table 1). AlShamaileh et al. [28] employed a planetary ball mill to grind different ratios of urea and kaolinite for 120 min at a rotational speed between 200 and 700 rpm. Contamination with impurities while milling, the agglomeration of particles, a wide range of particle size distribution, and heat generation are a few challenges associated with the mechanical attrition method (Table 1). Ultra-high-speed cutting was used by Liu et al. [17] to prepare the nano-composite as a coating material for NNFs. In this study, purified natural kaoline was treated with

an alkaline ether sulfate. Then, the mixture was commingled and cut at a high speed of 30,000 rpm for 10 min. This process enabled intercalation and gelation to take place. Rudmin et al. [18] activated glauconite for the preparation of a glauconite–diammonium phosphate NNF using a mill ring or planetary milling at 700 rpm for 10 min (Table 1).

3.1.3. Hydrothermal Synthesis

Hydrothermal synthesis is a bottom-up method used in the synthesis of nanoparticles. In this method, nanoparticles are formed through chemical reactions that occur under high-pressure and high-temperature conditions in an aqueous solution. It offers precise control over the size, shape, composition, and crystallinity of the resulting nanoparticles. The desired properties of the nanoparticles can be obtained by optimizing the reaction conditions, such as temperature, pressure, reaction time, and precursor concentrations. Bhardwaj et al. [30] employed hydrothermal synthesis for preparing two zeolites named SZC (synthesized zeolite clinoptilolite) and SZM (synthesized zeolite montmorillonite). In this method, tetraethoxy silane (C2H5O)4Si (silica source), aluminum nitrate (aluminum source), potassium (potassium source), and sodium hydroxide (alkali source) are reacted in a Teflon-lined stainless-steel pressure vessel, which was kept in a preheated (150 $^{\circ}$ C) oven for 72 h (Table 1).

3.1.4. Co-Precipitation Method

The co-precipitation method is a chemical method for the synthesis of nanoparticles in which multiple precursor materials react and form simultaneous precipitation. This involves with mixing of two or more precursor solutions, causing a chemical reaction that leads to the formation of nanoparticles. Nano-zeolite particles were synthesized as a carrier for the formulation of NNFs using the co-preparation method [32]. In this preparation, sodium silicate solution and ethylene glycol were heated and stirred in a homogenous mixture. Subsequently, AlSO₄ and NaOH were added dropwise while stirring at 50–60 °C. The resultant product formed a gray-colored nano-zeolite (Table 1).

3.2. Nanoparticle Modification Methods

The surface properties, composition, or structural changes to increase its desired characteristics are achieved through the modification of nanoparticles. This can be achieved through various methods including surface modification, chemical functionalization, and coating techniques. Several modification techniques, namely high-energy electron beam irradiation, ozone treatment, applying surfactants, and catalytic oxidation, are a few methods employed in the formulation of NNFs.

Although the nanonature of attapulgite is beneficial in the formulation of NNFs, the rod structures tend to aggregate with each other due to the high surface area and nanoeffect [16,26]. This necessitates modifications of attapulgite to improve the dispersion and retain its nanocarrier property. Toward this, Zhou, et al. [16] applied high-energy electron beam (HEEB) irradiation to natural attapulgite, which separated the rods from each other and increased the effective surface area (Figure 5). However, Cai, et al. [26] applied ozone (O₃) oxidation and hydrothermal processes to increase the dispersion of nanorods and increase –OH active sites on the surface. The authors reported that increased active sites might help to form a micro-nano network with urea (Table 1).

Zeolites exhibit high loading ability for anions, but they typically have a low capacity for loading cations onto their pores. However, this limitation can be overcome through structural modifications, which enhance their cation loading capacity. Methods such as applying surfactants and thermal treatments are commonly employed for these purposes [32]. In a study conducted by Milani et al. [33], it was observed that nano-ZnO particles exhibited a similar tendency for particle aggregation in water suspensions, forming clumps at the micrometer scale (Table 1). This aggregation process was found to be faster for nanoparticles compared to bulk particles, primarily due to Brownian motion and the uniform particle

size of nano-ZnO [33,42]. Additionally, the high ionic strength and low surface charge of ZnO also facilitate this aggregation process (Table 1).



Aggregated Attapulgite

Dispersed Attapulgite

Figure 5. A schematic diagram of aggregated attapulgite dispersed after irradiation.

4. Crop Responses for Nano-Nitrogen Fertilizers

Several studies have been conducted to investigate the responses of NNFs on different crops, considering different climate and soil conditions. These studies primarily examined agronomic performance, yield response, nitrogen uptake, and physiological changes. The following section summarizes the crop responses observed when applying NNFs.

4.1. Yield Responses

Ever growing population, limited arable land, and declining land productivity pose significant challenges to the agriculture sector. Henceforward, it is crucial to focus on increasing crop yield per unit application of nutrients [1]. Several studies showed that NNFs significantly increased crop yield compared to conventional N fertilizers. For example, Rudmin et al. [18] formulated new nano-ADP-glauconite fertilizers and tested them on oats. It was found that these fertilizers significantly (p < 0.05) increased the yield by 4.6% compared to non-fertilized treatment (Table 2).

The application rate of NNF influences the yield of crops. A greenhouse experiment conducted by Rop et al. [44] showed that a lower application rate (266 kg ha⁻¹) of NNFs significantly (p < 0.05) decreased maize yield by 91–191% compared to urea (the application rate was 532 and 1064 kg ha⁻¹). However, maize yield significantly (p < 0.05) increased the yield by 11% for NNFs compared to urea when a higher application rate (1064 kg ha⁻¹) was employed (Table 2). This observation was consistent with the yield of capsicum and kale in the same study [44]. A higher application rate of 1064 kg ha⁻¹ NNFs increased capsicum and kale yield by 14% and 18.6%, respectively, for NNFs compared to conventional fertilizers (Table 2). In a study, nano-Zno and vegetable oil-coated urea significantly (p < 0.05) increased the grain yield of wheat than non-fertilized wheat [45] (Table 2). Karoline–urea NNFs applied to rice increased the yield by 80% compared to single urea application [46].

Upadhyay et al. [47] reported that nano-urea foliar application with basal urea at low and high application levels did not significantly increase the yield compared to basal application of urea on maize and wheat in a two-year field study. However, 75% of the recommended level of basal application with foliar application of urea significantly increased (p < 0.05) maize yield in the last year of the study but was not effective for wheat in both years (Table 2). Similarly, the same treatment significantly (p < 0.05) increased the yield of mustard in the first year of study but not pearl millet [47]. Ammonium lignin-modified bentonite-coated urea formulations were prepared and tested on tomatos against urea in a greenhouse [22]. Most of the NNF formulations at different application levels provided significantly (p < 0.05) higher yields than urea (Table 2). However, when comparing a lower level of NNF (<100 kg N ha⁻¹) to conventional urea (100 kg N ha⁻¹), the NNF only yielded similar levels of crop yield. Raguraj et al. [48] tested urea–HA nanohybrid fertilizer against urea in tea fields across three agroecological regions in Sri Lanka (Table 2). According to the results, 10–17% and 14–16% yield increment was observed over urea in the low country

and Uva region, respectively, but there was no difference in the mid-country. Only a few studies compared the nano and bulk particles used to prepare SRFs on the crop responses. For instance, Dimkpa et al. [45] showed that coating urea with nano and bulk ZnO did not show any significant differences in the wheat yield (Table 2). In summary, studies have shown that NNFs can be a viable option to increase crop yield.

Table 2. The effect of N-nanofertilizers or synergists on crop response (significant values mean p < 0.05).

Fertilizer and Nanoparticle	Application Rate (kg/ha)	Crop	Crop Response	Study Country	References
Urea (U) + nanocarbon (NC) synergist	N—525 NC—1.575	Wheat	Leaf N accumulation is significantly higher by 55–65% than the control Glutamine synthetase activity and the nitrate transporter gene were higher than the control	China	[10]
Urea (U) + nanocalcium Carbonate (NCa) synergist	N—525 NCa—1.575	Wheat	Leaf N accumulation is significantly higher by 20–30% than the control Glutamine synthetase activity and nitrate transporter gene were higher than the control	China	[10]
Urea+ carboxylated nanocellulose	16.45	Wheat	Germination rate, tiller number, photosynthetic rate, and chlorophyll content were higher than urea treatment	China	[24]
Urea/NH4NO3 in attapulgite sodium polyacry-late polyacrylamide complex	81	Corn	Higher ¹⁵ N abundance and TN in the leaf Increased the height and stem diameter more than the control	China	[16]
Nano-nitrogen chelate (NNC) fertilizers	80–161	Sugarcane	The NUE of NNC was significantly higher than urea (control) treatment	Iran	[49]
Nano ADP—glauconite	50	Oat	The germination rate, plant height, and yield were significantly ($p < 0.05$) higher than the non-NNF-treated plot	Russia	[18]
Nano-hydroxyapatite (nHA) with cellulose fiber and polyacrylamide + urea	45–223	Maize	At a lower application rate, growth parameters were significantly ($p < 0.05$) lower than conventional fertilizers (CFs). However, at a high application rate, no significant difference was observed	Kenya	[44]
nHA with cellulose fiber and polyacrylamide + urea	45–223	Kale	At a high application level, NNFs showed significantly ($p < 0.05$) higher yields than CFs At a low application rate, herbage N was significantly lower by 33% than CFs	Kenya	[44]
nHA with cellulose fiber and polyacrylamide + urea	45–223	Capsicum	At a high application level, NNFs showed significantly (<i>p</i> < 0.05) higher yields by 54% than conventional fertilizers At a low application rate, herbage N was significantly lower by 43% than CFs	Kenya	[44]
Zno-np/vegetable oil (VO)-coated urea (ZN-VO-urea)	100 mg-N kg ⁻¹ soil	Wheat	ZN-VO-urea fertilizers showed significantly (<i>p</i> < 0.05) higher yields than VO-coated urea. But, plant N was not significantly different between them However, there was no significant difference between nano- and bulk-ZN-VO coated urea. This suggests that Zno has a greater synergetic effect with fertilizer than the nanosize of the particle in	United States	[45]
Nano-urea	0–150	Maize and mustard	At a 113 kg-N ha ⁻¹ application rate, nanofertilizers showed significant ($p < 0.05$) yields compared to CFs	India	[47]
Quaternary ammonium lignin (QAL)-modified nano-bentonite-	75–300	Tomato	Most of the NNF significantly increased the yield and N uptake by tomatoes than urea	Egypt	[22]
coated urea Urea–HA nanohybrid fertilizer	240	Tea	A tea yield increase was noticed in the low country and Uva region but not in mid-country	Sri Lanka	[48]
Urea-chitosan nanohybrid fertilizer (UCNH)	Urea (66–165 kg N ha ⁻¹) + urea-chitosan (0–500 mg N L ⁻¹)	Rice	The best treatment was the application of a 500 mg N L^{-1} compensatory level of UCNH with 60% of the recommended urea level (99 kg N ha ⁻¹)	Egypt	[50]

Fertilizer and Application Rate Nanoparticle (kg/ha)		Crop	Crop Response	Study Country	References
Urea surface-modified hydroxy appetite (HA) nanoparticles	0–33 kg N ha $^{-1}$	Almond	Nanofertilizer at a higher application rate significantly increased the germination of almonds more than urea and ammonium sulfate	Egypt	[51]
NNF	Ammonium nitrate (AN) (0–100%) and/or NNF (0–75%)	Lettuce	In both study years, NUE significantly increased for a 100% NNF application than a 100% AN application	Egypt	[52]
Kao-urea NNF	150 kg ha^{-1}	Rice	The best NNF significantly increased the yield but not the leaf N content compared to urea	Malaysia	[46]

Table 2. Cont.

4.2. Crop Nitrogen Uptake

The uptake of nitrogen by crops is vital for promoting healthy plant growth, protein synthesis, and photosynthesis, ultimately leading to increased crop productivity. Efficient nitrogen uptake also contributes to environmental sustainability by minimizing nutrient waste and reducing the environmental impact of agricultural practices. Studies showed mixed effects of NNFs on crop nitrogen uptake. For instance, ¹⁵N labeled urea in an attapulgite sodium polyacrylate polyacrylamide complex was tested against urea alone on corn in a pot experiment [16]. This NNF showed significantly (p < 0.05) higher plant total ¹⁵N compared to urea (Table 2). Rop et al. [44] developed a cellulose-graftpoly(acrylamide)/nanohydroxyapatite NNF and tested it on maize, kale, and capsicum. At a 1064 kg ha⁻¹ application rate, NNF significantly (p < 0.05) lowered capsicum herbage N compared to urea. However, maize and kale herbage N were not significantly different between NNF and urea treatment (Table 2).

Dimkpa et al. [45] tested nano-ZnO/vegetable oil (VO)-coated urea fertilizer against urea with nano-Zno (without coating) on wheat under drought and non-draught conditions (Table 2). The results revealed that both fertilizers gave similar plant N uptake under drought and non-draught conditions. In another study, Helal et al. [22] examined ammonium lignin-modified bentonite-coated urea formulations that were applied to tomatoes in a greenhouse study. These formulations significantly (p < 0.05) increased the N accumulation in plants than urea (control). The highest N accumulation was 4.7 times higher than the control (Table 2). A study tested developed urea-chitosan nanohydride fertilizers (UCNH) and tested them on rice [50]. Different levels of UCNH (0–500 mg N L^{-1}) were applied as compensation for urea along with conventional urea (165–66 kg N ha⁻¹), and N uptake by grains and straw was measured. When the UCNH percentage increased and the urea level decreased, the N content of both cereal and straw significantly (p < 0.05) decreased. Nevertheless, a 500 mg N L⁻¹ compensatory level of UCNH with 60% of the recommended urea level (99 kg N ha⁻¹) did not significantly (p < 0.05) decrease the N content, and is thus suggested as the best alternative for 100% recommended urea level application (165 kg N ha⁻¹) (Table 2). An NNF made of urea and kaolinite also has been reported to not increase the leaf N compared to urea application [46].

Nanomaterials are used as synergists with conventional fertilizers to increase crop performances in sustainable agriculture [53]. A study used nono-carbon, nano-CaCO₃, and the composite of both as synergists with urea on winter wheat to investigate their performance in pot and field experiments [10]. The results indicated that all the synergists had a significant (p < 0.05) effect on increasing plant nitrogen (N) uptake, with improvements ranging between 15% and 65% compared to using urea alone (Table 2). Another study reported that the nitrogen uptake of peanuts increased with the co-application of nano-CaCO₃ [14] (Table 2). Nanomaterials increase the dispersion of fertilizers in the soil [54] and also act as an agent to control nutrient release [10,53]. Ammonium ions are adsorbed on the surface of nanomaterials and release H ions, which control the loss of ammonium ions in soil [10]. For instance, non-conductive nanocarbon absorbs the N of a NH₄⁺ ion and releases a H⁺ ion. Further, nanomaterials inhibit denitrifying bacteria, thus

decreasing nitrate losses [55]. As a result, the reduction in nutrient loss leads to an increase in plant uptake.

4.3. Nitrogen Utilization Efficiency (NUE)

Nitrogen utilization efficiency (NUE) refers to the ability of crops to convert the applied nitrogen to a useful component (i.e., yield). Improving NUE is essential for sustainable agriculture, as it helps minimize N losses, thus reducing environmental pollution and optimizing crop productivity. Only a limited number of studies have reported the NUE of NNFs. For example, a study reported that the NUE of winter wheat increased by 5–22% by applying nano-carbon and nano-CaCO₃ as synergists [10] (Table 2). An HA-urea nanohybrid applied to tea in Sri Lanka significantly (p < 0.05) improved NUE [48] (Table 2). In a field study, an NNF was applied to lettuce as a foliar application at different levels, with ammonium nitrate (AN) as a compensator for AN [52]. NUE was significantly (p < 0.05) three times higher for a 100% NNF application than a 100% AN application in both years (Table 2). Some other studies suggested that this could be due to the regulation of important nitrogen metabolism-related unigenes [56].

4.4. Germination of Seeds

Few studies have reported that NNFs contribute to an increase in germination percentage in many plant species. A study reported that a urea/carboxylated nanocellulose NNF increased the germination percentage of wheat by 20–27% compared to urea [24] (Table 2). Nano-ADP-glauconite, an NNF, significantly (p < 0.05) increased the germination percentage of oats by 6% for the best treatment [18] (Table 2). Badran et al. [51] conducted a comprehensive study to examine the influence of urea, ammonium sulfate, and NNFs (urea surface-modified hydroxy appetite nanoparticles) on the germination of almond seeds under varying levels of saline conditions. The study found that the germination rate was significantly (p < 0.05) higher for NNFs compared to other fertilizers at all application rates and under all levels of saline conditions (1, 3, and 5 ds m⁻¹) (Table 2). However, germination percentage significantly (p < 0.05) improved only at higher application levels of NNFs under all saline conditions. Therefore, this study shows that NNFs can be used to alleviate the effect of saline water on seed germination. The higher germination exhibited by NNFs could potentially be attributed to the intake of nanomaterials by seeds, leading to an increase in the micropores for water intrusion that could increase germination [57].

4.5. Chlorophyll Content

Chlorophyll is a color pigment that is responsible for capturing light energy during photosynthesis, and its content directly influences energy production and essential metabolic processes. Studies have shown that the application of NNFs enhances chlorophyll content in plants compared to conventional nitrogen fertilizers. For example, a 25% AN and 50% NNF application significantly (p < 0.05) increased the chlorophyll content of lettuce by 10 SPAD compared to AN application alone in two years [52]. This observation is directly correlated with the leaf N content. Quaternary ammonium lignin (QAL)-modified nano-bentonite-coated urea applied to tomatoes significantly (p < 0.05) increased the leaf chlorophyll content compared to urea at higher application rates [22]. However, at lower-level applications, the chlorophyll content is only comparable to urea.

4.6. Gene Expression

Several studies reported the positive influence of NPs on gene expression [58–61]. However, only a few studies reported that NNFs induce gene expression, which is beneficial for crops. Glutamine synthase (GS) genes are responsible for N assimilation in crops, which induce nitrate transportation family genes. Yang et al. [10] found that nanocalcium carbonate (NCa) and nano-carbon (NC) synergists applied to wheat with N fertilizers significantly (p < 0.05) increased the expression of GS, such as TaGS1, TaGS2, TaNRT2.2, and TaNRT2.3, compared to N fertilizer alone. Therefore, these synergists increased the N

transportation and accumulation in wheat. Chew et al. [62] reported that biochar-based nano-iron-fortified compound fertilizer significantly (p < 0.05) downregulated the expression of ammonium transportation genes such as OsAMT1.1, OsAMT1.2, and OsAMT1.3 in roots compared to the control, whereas this NNF significantly (p < 0.05) increased nitrate transporter genes, such as OsNAR2.1 and OsNRT2.3, in roots compared to the control. It is evident that an NNF promotes the transportation of nitrates over ammonium ions.

5. Disadvantages of Nanoparticles for Crops and the Environment

Despite nanoparticles providing several benefits to crops and the environment, a few disadvantages were also identified that are associated with them. Excess application of nanoparticles could be toxic for plants. The toxicity of the NPs depends on their physiochemical properties such as particle size, reactivity, charge density, shape, and solubility [63]. Dimkpa et al. [45] reported that nano-ZnO can be toxic to plants and, therefore, NNFs should formulated with low concentrations of nano-ZnO. Nevertheless, there is currently no direct evidence demonstrating the toxicity of NNFs for plants.

Several studies showed that NPs are ingested by plants and modify biological activities such as nutrient uptake, photosynthesis, and gene expression [64]. In a similar way, NPs can mutate plant genes and induce them to produce chemical compounds that are toxic and detrimental to the plants themselves and consumers. Additionally, these NPs can react with biochemicals within the cell and form unwanted chemical compounds.

NPs have been slowly accumulating in the environment and can potentially harm the soil, crops, and ecosystem. Metal and ion NPs used for the formulation of NNFs could persist in the soil for a long time, which may be toxic for crops and may change the nutrient dynamics. NPs with heavy metals contaminate the soil and water bodies. Some of the NPs possess antimicrobial activity, which can adversely impact the beneficial soil microbial population [65]. Nevertheless, no comprehensive studies were conducted to examine the impact of residual NPs in soil and the fate of heavy metals.

The application of dry nanoscale fertilizers through broadcast methods can result in drift losses, leading to adverse consequences for human respiratory health [45]. Several studies highlighted the negative impact of NPs on human cells like damage to the plasma membranes, toxicity for mitochondria, nuclear damage, the formation of reactive oxygen species, and interference with signaling pathways [66].

Currently, relevant regulatory frameworks to deal with nano-based products, including NNFs, are unavailable. The leading countries producing nano-based products are developing safety standards, risk assessment protocols, and regulatory frameworks to safeguard the workers in the relevant industries and the consumers [66].

6. Limitations in the Studies and Future Research Directions

- It is worth noting that certain studies have compared nanofertilizers (NNFs) to untreated control groups, which inherently results in superior performance by the NNFs. However, to provide a more comprehensive evaluation, it is important for studies to include appropriate control groups for comparison. These control groups should consist of conventional fertilizers, other smart fertilizers, or other commercially available nanofertilizers commonly used in agricultural systems. Comparing NNFs against these control groups allows for a more accurate assessment of the effectiveness and added benefits of newly developed NNFs.
- Only a limited number of studies have directly compared the effects of real nanoparticles with regular particle sizes on crop responses [31,33,45]. This comparison is crucial to understand the unique impacts of nanoparticles and to differentiate them from the effects of conventional particle sizes. Therefore, future studies need to focus on comparing the effect of nanoparticles on controlling nutrient release.
- Many studies on nanoparticles have been conducted in controlled laboratory settings, which may not fully represent real-world agricultural conditions. More field

studies are required to validate the findings and assess the practical implications in agricultural settings.

- Nanoparticles have the potential to persist in the environment over the long term. It is
 important to understand the fate and behavior of these particles to assess any potential
 risks. However, currently, there is a lack of publicly available studies specifically
 addressing long-term trials and their impact on the environment. Further research is
 needed to comprehensively evaluate the long-term effects of nanoparticles and ensure
 their safe and sustainable use in agricultural applications.
- Some nanocarriers have the potential to be phytotoxic, which may have negative impacts, namely stunted growth, reduced biomass accumulation, chlorosis, wilting, and even plant death. Therefore, studies focusing on the accumulation of nanoparticles by plants and their long-term effects need to be conducted.
- Further research is needed to investigate the transmission of nanoparticles (NPs) through the food chain. Understanding how NPs can potentially accumulate and transfer from one organism to another within the food web is crucial for assessing their overall impact on human health and the environment.
- The fate of NPs ingested by the crops and their negative impacts on the crop itself and the living beings consuming them needs to be analyzed rigorously. This will increase the understanding of the current gray area of nanoparticle (NP) behavior and its potential risks in the food chain.

7. Conclusions

This review article provides a comprehensive summary of the existing literature concerning the formulation of nano-nitrogen fertilizers and the corresponding crop responses upon their application with a special emphasis on their contributions to the Sustainable Development Goals (SDGs). Studies demonstrated that using NNFs provided benefits over conventional N fertilizers with respect to yield, quality of yield, nitrogen utilization efficiency, and various physiological responses. Therefore, NNFs have a high potential to address SDGs, such as alleviating hunger and mitigating environmental contamination with N fertilizers. However, there are gray sides to NNF applications since the negative impacts of these fertilizers on human health, soil microbial population, other flora and fauna, and the total ecosystem are not completely understood yet. Considering these factors, it is imperative to establish regulations for the application of NNFs and develop global standards for evaluating their impacts.

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