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Zein as a renewable material for the preparation of green nanoparticles for drug delivery

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Environmental sustainability is a key challenge driven by the increased consumption of natural resources with limited availability. In this scenario agriculture has emerged as a privileged source of renewable resources, hence more efforts should be addressed to the study of plant-derived materials for medical applications. Zein is a biocompatible, biodegradable and amphiphilic prolamin protein extracted from the endosperm tissue of corn. For these reasons, its applications span from coatings for edible capsules, to the fabrication of bi- and tridimensional scaffolds for tissue engineering, and to develop drug delivery systems. This review aims at describing the properties and main applications of zein with a focus on the most recent and updated state of the art literature on zein based nanoparticles for the controlled delivery of various drugs. The main focus is to analyze the state of the art literature to understand how to implement sustainable methods for the preparation of zein NPs and to propose their exploitation as novel drug delivery systems for multiple applications, including oligonucleotide delivery. Main methods for zein NP preparation are described under an ecofriendly point of view, highlighting their environmental sustainability based on used solvents, waste products and energy consumption.

KEYWORDS

drug delivery, green technologies, nanoparticles, zein, sustainable materials

1 Introduction

The continuous growth of the world population (that according to predictions will reach around 8.5 billion in 2030 and 9.7 billion in 2050) has tremendously increased the demand for natural resources, which however show limited availability (Pimentel and Pimentel, 2006; FAO, 2013; Gaigbe-Togbe et al., 2022). Based on that, new environmentally-sustainable materials and technologies should be urgently developed (Chakrapani et al., 2022; Tamasiga et al., 2022), following the “12 Principles of Green Chemistry” (Anastas and Eghbali, 2010; Duarte et al., 2014) and the “12 Principles of Green Engineering” (Brennecke, 2004) (Table 1). In this scenario, one main target is the shift from linear to circular economy, a new model of production and consumption based on the recycle and reuse of waste, aimed at the preservation of ecosystems and biodiversity. Based on the circular economy model, produced waste should be converted into renewable resources leading to the elimination of the concept of “waste” itself (Zimmerman et al., 2020).

Waste materials from agriculture (i.e., plant-derived materials) are particularly promising as green resources due to their high availability, renewability, low-cost, and

TABLE 1 The 12 principles of green chemistry (Anastas and Eghbali, 2010) and engineering (Brennecke, 2004).

| 12 principles of green chemistry | 12 principles of green engineering |
|--|---|
| Prevent waste | Minimize hazardous material and energy |
| Maximize atom economy | Prevent waste |
| Minimize hazardous chemical synthesis | Minimize separation and purification operations |
| Use safer solvents and auxiliaries | Maximize mass, energy, space, and time efficiency |
| Design energy efficiency | Design output-pulled rather than input-pushed |
| Use renewable feedstock | Design recycle and reuse based on complexity |
| Minimize or avoid derivatives | Target durability rather than immortality |
| Use catalysts, rather than stoichiometric reagents | Meet needs, minimize excess |
| Design safe degradation | Minimize material diversity |
| Analyze in real-time for pollution prevention | Integrate local material and energy flows |
| Minimize the potential accidents | Design for commercial “after-life” |
| Design safe chemistry for accident prevention | Look for renewable rather than depleting |

potential exploitation in the food industry (for example, zein from corn crop waste as edible carriers for flavor compounds, to deliver food colorants or to give luster to the coating materials in confectionery products) (Kasaai, 2018), as well as for biomedical and pharmaceutical applications (for example, pectin from juice, apple, and cider waste industries or starch from cereal and potato waste for bioprinting applications) (Jovic et al., 2019; Chakrapani et al., 2022). Plant-derived materials include: proteins (such as wheat gluten, soy and zein), polysaccharides (such as pectin, cellulose and starch), lignin and plant extracts rich in polyphenols (such as Aloe Vera) (Iravani and Varma, 2019). Among them, plant-derived proteins show promising properties for applications in drug delivery and tissue engineering (Akharume et al., 2021; Tortorella et al., 2021). Cereals by-products provide a valuable source of prolamins, which are the main storage proteins present in most cereals (Song et al., 2021). Among them, herein we focus on zein, which is the major storage protein present in the cytoplasm of endosperm cells of maize (*Zea mays*), constituting almost 50% of corn proteins (Corradini et al., 2014). Other cereals, such as wheat, barley and sorghum are sources for other plant-derived proteins, respectively gliadin, hordein and karifin, with similar properties to zein. However, compared to other plant-derived proteins, zein is advantageous due to the high availability of materials for its extraction (from agricultural waste) and high extraction yield (Anderson and Lamsal, 2011). Indeed, corn is the main global staple cereal in terms of annual production, exceeding 1 billion metric tons and expected to rise with an estimated annual increase of 37 million tons (Erenstein et al., 2022). Feedstock for zein extraction can be obtained from a primary process (including agricultural crops) or from a secondary process (which includes agro-industrial waste resulting from the processing of maize as a by-product by wet-milling or dry-milling) (Tapia-Hernández et al., 2019). Prolamin extraction is generally carried out by washing with aqueous ethanol; however, sustainable and innovative protocols for zein extraction have recently been developed to minimize environmental impact of prolamin processing, reducing energy consumption and solvent

volumes (Chen et al., 2022a). Zein is also characterized by other favorable properties, such as biocompatibility, biodegradability and amphiphilicity, which is responsible for its interaction with both hydrophobic and hydrophilic materials and/or drugs. Zein has also been certified as Generally Recognized As Safe (GRAS) in the 1985 and approved as safe biomaterial for medical use by the Food and Drug Administration (FDA) (Weissmueller et al., 2016; Labib, 2018). Previous literature has reported the use of zein for the production of edible capsules (Bouman et al., 2016), films (Wang et al., 2005; Zhang et al., 2015; Hasanzadeh et al., 2021), hydrogels (Gao et al., 2007), scaffolds for tissue engineering (Torres-Giner et al., 2009; Ali et al., 2014; Khatri et al., 2020) and drug-loaded NPs (Abdelsalam et al., 2021). Due to its promising properties, zein has been the subject of previous review papers, focused on zein extraction and general applications (Shukla and Cheryan, 2001; Anderson and Lamsal, 2011), as well as zein biomedical applications (Corradini et al., 2014; Paliwal and Palakurthi, 2014; Pérez-Guzmán and Castro-Muñoz, 2020; Tortorella et al., 2021) including the design of zein-based NPs (Zhang et al., 2016; Abdelsalam et al., 2021; Yuan et al., 2022a; Yuan et al., 2022b). Zein NPs are advantageous over lipid and synthetic polymer NPs due to their superior biocompatibility, while compared to NP based on animal-derived proteins, they are not affected by the risk for zoonotic disease transmission (Reddy and Yang, 2011; Podaralla and Perumal, 2012). Furthermore zein is amphiphilic, as composed by hydrophobic and neutral amino acids, such as leucine, proline and alanine, and by hydrophilic glutamine residues (Shukla and Cheryan, 2001). Zein amphiphilicity affects its solubility and drug loading ability: 1) zein is insoluble in water and soluble in water-alcohol, ketone and highly concentrated salt and alkaline solutions; 2) zein NPs can encapsulate both hydrophobic and hydrophilic drugs. However, in most studies zein NPs have been used for the release of food and nutrient components (Kasaai, 2018), and lipophilic drugs, such as curcumin (Shukla and Cheryan, 2001), gallic acid (Wiggers et al., 2022), gitoxin (Muthuselvi and Dhathathreyan, 2006) and fluorouracil (Lai and Guo, 2011). Only

a few recent reports have proposed the encapsulation of hydrophilic drugs (e.g., doxorubicin and chloroquine) (Dong et al., 2016; Chen X. et al., 2022) or plasmid DNA (El Sharkawi et al., 2017) into zein NPs. Such recent studies exploit the possibility to vary zein charge depending on the pH to promote its electrostatic interactions with hydrophilic molecules with opposite charge, enhancing drug encapsulation efficiency. Furthermore, surface coatings have been applied on zein NPs to prepare core-shell nanocarriers with specific properties, including enhanced stability to centrifugation and freeze-drying (Tortorella et al., 2021). As zein has the ability to protect labile drugs from degradation and is more resistant to degradation than other proteins, it has been widely applied for oral drug delivery (Luo et al., 2010; Lee et al., 2013). However other routes of administration, such as nasal, transdermal delivery, intramuscular, intravenous and subcutaneous delivery have also been exploited (Hurtado-López and Murdan, 2010; Lai and Guo, 2011; Li H. et al., 2019; Yu et al., 2022).

This review provides an initial overview on the chemistry and structure of zein molecules as preliminary to a focus on the use of zein as a green material in the field of nanomedicine, exploiting sustainable preparation processes. Hence, novelty of this review derives from focusing on zein as a byproduct material from agricultural waste, with proper physicochemical properties for the preparation of added-value pharmaceutical products under environmentally friendly production processes. To this aim, methods to prepare zein NPs are initially described and their sustainability is discussed, in terms of solvent types and ratio, consumed energy and produced waste. The possibility to exploit zein NPs for the loading and release of hydrophilic drugs, such as oligonucleotides, is also discussed, taking advantage of the superior biocompatibility of zein over state of the art materials for nanocarrier production. Potential issues associated with the *in vivo* administration of zein NPs, such as immune response, are also critically discussed (Zhang et al., 2016).

Extensive state of the art research has been carried out, covering a wide range of publications with a focus between 2017 and 2022 to highlight the most innovative preparation processes and applications of zein in the drug delivery field. However, older literature was also considered for the analysis of zein physicochemical characteristics. The main databases employed for state of the art research were Science Direct, PubMed and Scopus with a focus on keywords such as “zein nanoparticles” and “green drug delivery systems”. The innovative contribution of this review paper is to inspire the future application of zein derived from agricultural waste for the preparation of green NPs by sustainable methods, and to encourage their use in advanced therapies, such as the release of RNA molecules.

2 Chemical and molecular structure of zein

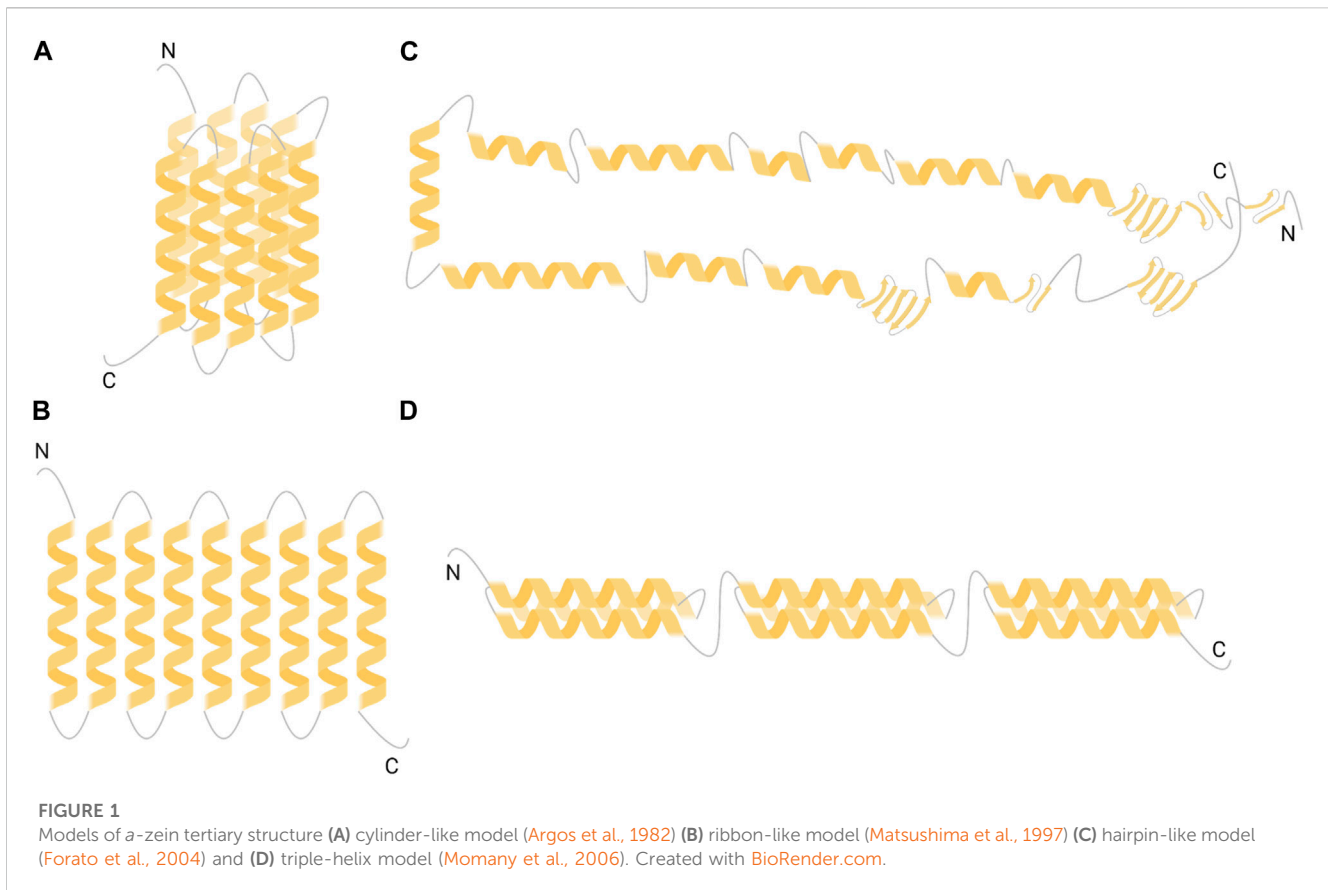
2.1 Composition of plant-derived zein

Zein has a peculiar aminoacidic composition, with a prevalent proportion of nonpolar amino acids respect to polar ones. Hydrophobic and neutral amino acid residues include a high percentage of leucine (~20%), proline (~10%) and alanine

(~10%), while hydrophilic residues have a content of approximately 21%–26% glutamic acid (Abdelsalam et al., 2021). Notably, the following amino acids are present at very low amounts: lysine (~0%), tryptophan (~0.16%), arginine (~5%) and histidine (~1%) (Geraghty et al., 1981; Shukla and Cheryan, 2001). Overall, zein is an amphiphilic protein with an isoelectric point of 6.2–6.8, close to the physiological pH. Therefore, at physiological pH zein aggregation occurs due to enhanced hydrophilic interactions and decreased electrostatic repulsions. On the other hand, at low and high pH zein is charged: at acidic pH it acquires an overall positive charge while at basic pH it becomes negatively charged. Therefore, zein NPs are often exploited for oral delivery of compounds to prevent their degradation in the harsh gastrointestinal tract (GIT) conditions because at acidic pH they remain stable against aggregation. Zein NPs are often studied in simulated GIT fluids (at acidic pH, with pepsin) to evaluate NPs stability (Abdelsalam et al., 2021). The functionality of zein is given by its numerous functional groups, among them amines, amides, hydroxyls, carboxylates and phenols (Corradini et al., 2014). Amines from glutamine residues are more exposed on the peptide chains thanks to the tertiary structure of zein, described in the Par. 2.2. Crosslinking of zein could be obtained using chemical crosslinking agents, such as EDC/N-hydroxysuccinimide (NHS), formaldehyde, glutaraldehyde, succinic acid, citric acid and eugenol (Corradini et al., 2014; Tortorella et al., 2021). The self-crosslinking of zein molecules mediated by EDC/NHS was reported for film fabrication, linking amines with carboxyl groups (Kim et al., 2004). Another study describes zein films prepared by using succinic anhydride, succinic acid, citric acid, or eugenol, separately or together, as natural crosslinkers to increase their tensile strength (Khalil et al., 2015). Zein electrospun membranes were crosslinked through non-toxic citric acid, thus forming amido linkages between the carboxyl groups of citric acid and the amino groups of zein. Results showed higher mechanical properties, longer water stability and higher cell adhesion compared to uncrosslinked electrospun nanofibers (Jiang et al., 2010). Zein based crosslinked hydrogels were also prepared with increased absorption capacity, using N,N'-methylene bisacrylamide as the crosslinker (Ni et al., 2018), or with increased mechanical properties combining chemical (through glutaraldehyde, linking amine and aldehyde groups) and physical crosslinking (through hydrophobic interactions and hydrogen bonding upon a post-treatment with water) (Wang et al., 2019).

Composition confers zein unique solubility behavior: zein is insoluble in pure water, while it is soluble in water-alcohol, water-acetone, highly alkaline and urea solutions, and in the presence of anionic detergents (Lawton, 2002; Deo et al., 2003; Paliwal and Palakurthi, 2014). Due to the abundance of nonpolar amino acids and the insolubility in water, zein has been initially investigated for the preparation of tough greaseproof coatings of films resistant to acid and microbial attack, adhesives for the introduction of flavors and medical tablets (Shukla and Cheryan, 2001; Liu et al., 2005; Zhong and Jin, 2009). Only recently, zein has been proposed for the preparation of NPs for controlled and sustained drug release (Zhang et al., 2016; Tran et al., 2019; Abdelsalam et al., 2021).

Zein can be classified into four main groups with different peptide composition, molecular weight, solubility and charge: α -zein, β -zein, γ -zein and δ -zein. Zein contains a mixture of them and has an average molecular weight of 10–30 kDa. The



main component is *a*-zein, representing around 80% of the total prolamin content present in corn. This protein fraction has an average molecular weight of 19–22 kDa and is the most abundant in commercial zein, among the four groups (Cabra et al., 2006; Cabra et al., 2007). The second most abundant fraction is β -zein, which represents 10%–15% of zein and has an average molecular weight of 17–18 kDa. Respect to the aminoacidic composition of *a*-zein, β -zein has less proline, arginine, methionine and histidine. γ -zein shows the highest molecular weight (27–28 kDa), but it constitutes only 5%–10% of zein. Finally, δ -zein is the minor fraction in zein (only 3%) and has the lowest molecular weight (9–10 kDa) (Esen, 1986; Wilson, 1991; Parris and Dickey, 2001). Among major food and pharmaceutical companies, Flo Chemical Corp (Ashburnham, MA), which is specialized in distribution of zein for various applications, and Showa Sangyo (Tokyo, Japan), that sales zein together with other food derivatives, hold the record in zein commercialization (Shukla and Cheryan, 2001; Anderson and Lamsal, 2011).

2.2 From composition to structure of zein protein molecules

Literature has reported various studies aimed at elucidating the tertiary structure of *a*-zein, being the mostly available zein fraction (Argos et al., 1982; Garratt et al., 1993; Matsushima et al., 1997; Forato et al., 2004; Momany et al., 2006). Molecular models of *a*-zein tertiary structure include cylinder model, ribbon-like model, hairpin

model and super-helical structural model (Figure 1). Such models approximate *a*-zein as composed by two main fractions with 19 kDa and 22 kDa molecular weight (referred to as Z19 and Z22, respectively), based on data on zein composition obtained from sodium dodecyl sulfate-polyacrylamide gel electrophoresis analysis (SDS-PAGE) (Lee et al., 1976).

The cylinder model is one of the firstly developed structural models of *a*-zein (Figure 1A) and was derived from circular dichroism (CD) data (Argos et al., 1982). CD spectrum of a mixture of Z19 and Z22 fractions in 70% methanol aqueous solution was measured in the 190–240 nm wavelength range. Data suggested that *a*-zein tertiary structure prevalently consists of α -helices (50%–60%), with turns or random coils in the remaining chain portions. Notably, α -helix content was in agreement with data from previous optical rotatory dispersion measurements, performed on zein solutions in 50%–80% ethanol-water (Danzer et al., 1975). According to the cylinder model, 9 homologous repeating α -helix units are topologically arranged adjacent and antiparallel, forming a cylindrical structure with turns linking the α -helices (Figure 1A). The cylinder has an oval cross-section with the helix and cylinder axes aligned. As α -helices have mainly a hydrophobic nature, with only a few polar amino acids, the cylinder-like structure shows hydrophobic behavior except for the top and bottom surfaces, containing turns rich in hydrophilic glutamine residues. This cylindrical structure with polar glutamine-rich turns is stabilized by side chain interactions, specifically Van der Waals interactions and hydrogen bonds.

The ribbon-like model (Figure 1B) was developed based on data from small-angle X-ray scattering (SAXS) analysis, performed on

a-zeins solutions in 70% v/v ethanol-water. Data suggested the presence of 9 and 10 repeating *a*-helix units in Z19 and Z22 respectively, joined by glutamine-rich loops (Tatham et al., 1993). The ribbon-model described by Matsushima et al. is based on the linear stacking of 9–10 adjacent antiparallel *a*-helices, perpendicular respect to the helical axis, and does not contain β -sheet, forming an elongated prism-like structure with $13 \times 1.2 \times 3 \text{ nm}^3$ size (Matsushima et al., 1997) (Figure 1B). Therefore, this two-dimensional ribbon-like structure is organized with hydrophilic ends and hydrophobic sides, enhancing the stability of *a*-zein. This slender prism model with hydrophilic extremities and hydrophobic surfaces was also employed to understand the orientations of zein molecules on hydrophilic and hydrophobic surfaces, as well as zein self-assembly from single molecules into NPs (Wang et al., 2008; Wang and Padua, 2012).

The hairpin-like model was developed, based on nuclear magnetic resonance (NMR), SAXS and Fourier transform infrared spectroscopy (FTIR) analyses, performed on Z19 fraction only (Forato et al., 2004). Particularly, FTIR analysis showed that Z19 structure contains a high content of *a*-helices (46%) and a limited amount of β -sheet (22%). In the hairpin-like model, helices are connected to each other to form rings, sheets or loops, with dimensions coherent with the previous model proposed by Matsushima and coworkers (Figure 1C). This structure has the ability to refold on itself or to extend based on the employed solvent. The hairpin-like model confirmed the presence of 20 amino acids for each *a*-helix, as stated in previous models (Argos et al., 1982; Matsushima et al., 1997). The hairpin-like model is useful to explain the process of zein fiber formation.

More recently, a super-helical structural (or triple-helix) model of Z19 tertiary structure was developed, based on molecular mechanics and dynamic simulations, in order to overcome incoherencies between previous models and experimental data (Momany et al., 2006). In this model, each super-helix contains three antiparallel *a*-helices, linked together by glutamine-rich turns (Figure 1D). Probability algorithms suggested that *a*-zein mainly forms coiled coils rather than pure *a*-helices. Coiled coils show a slight curvature along the helical axis, conferring the helices a twist spatial organization. Based on the amino acid composition of a single *a*-helix (from 14 to 19 residues), the *a*-helix percentage was estimated to be 62%, which is consistent with FTIR data.

3 Zein micro and nanoparticles: towards green methods for their production

3.1 Solvent selection for zein solubilization

Solvent selection is important for zein extraction, purification and for the manufacturing of zein substrates, including products for regenerative medicine, such as scaffolds, hydrogels or particles. Generally, zein can be solubilized into three types of solvents: primary, secondary and tertiary solvents. Primary solvents are single solvents able to dissolve zein such as glycerol and acetic acid (Evans and Manley, 1941). They generally have hydroxyl, amine, amide and carboxyl functional groups. Indeed, solvents with the above characteristics display balanced polar and

nonpolar features and are thus able to dissolve amphiphilic zein. Typically, primary solvents produce more stable zein solutions than binary solvents containing additional water (Evans and Manley, 1941; Anderson and Lamsal, 2011). On the other hand, studies on zein solubility showed its improvement in binary solvent mixtures by introducing water (Dill, 1927; Swallen, 1941). Different binary solvent combinations were studied and results showed that water-anhydrous alcohol, water-ketone, aromatic hydrocarbon-anhydrous alcohol, and alkaline agent-water represent effective secondary solvents for zein. For each secondary solvent, solubility varies depending on the composition of the binary mixture and on other parameters, such as temperature. An in-depth understanding of zein solubility in secondary solvent mixtures has required throughout investigations (Manley and Evans, 1943). In detail, experiments were carried out by adding minor amounts of aliphatic alcohols, ketones, or glycols to water, aromatic hydrocarbons, chlorinated hydrocarbons, nitroparaffins, aldehydes, or cyclic ethers. Results showed that zein solutions in aqueous binary mixtures show time and heat dependent behavior, with possible undesirable formation of gels or foams respect to non-aqueous systems with increasing zein and water concentration. In addition to widely used alcohol-water solvents, mixtures of acetone, acetonyl-acetone, dioxane, and dioxolane with water are also good solvents for zein. Particularly, zein solutions in acetone-water solvents have shown lower viscosities respect to solutions in other binary solvents with the same concentration. Indeed, zein solutions with 20% w/v concentration in 70% v/v acetone-water and in 70% v/v isopropyl alcohol-water showed viscosity of 20 cP and 83 cP at 30°C, respectively. Solution viscosity is an important parameter for the selection of the most appropriate solvent mixture for zein solubilization. Solution viscosity value is also relevant to assess zein absorption rate from solutions, to obtain zein coatings on substrates, such as films and NPs.

Finally, a thorough investigation of ternary solvents for zein was also conducted using both aqueous and non-aqueous mixtures (Evans and Manley, 1944). Such studies established that zein solutions in aqueous ternary solvents containing water, acetone and an aldehyde showed similar behavior to zein solutions in aqueous alcohol and aqueous aldehyde binary mixtures, but had higher resistance to gelling when at least 5% v/v aldehyde was added to the mixture because aldehydes can be employed as gelation inhibitors in aqueous zein solutions.

Despite the many studies on zein solution behavior as a function of different solvents, the effects of zein solution concentration, solvent ratio and temperature on zein conformation are still poorly known. More recent studies have focused on the understanding of the properties of zein products obtained from solutions in commonly used ethanol-water and acetic acid-water solvents (Fu and Weller, 1999; Shi et al., 2009; Zhong and Ikeda, 2012). Such research studies have found that zein films cast from ethanol-water or acetic acid-water solutions on silicon wafers have different surface morphologies and hydrophilicity, as assessed by tapping mode atomic force microscopy (TP-AFM), water contact angle measurements and X-ray photoelectron spectroscopy (XPS). In detail, zein films prepared from acetic acid-water solutions showed smoother surface and higher hydrophobicity respect to those prepared from ethanol-water solution. This behavior was explained based on the influence of acidic or basic environment

on the properties of zein molecules (Zhang et al., 2011). Indeed, in acidic solution (i.e., in acetic acid-water solutions), zein molecules tend to preferentially expose their hydrophobic groups in contact with air because the silicon wafer surface contains mainly silanol groups that confer a negative charge. On the contrary, zein films prepared from ethanol-water solutions at neutral pH are more hydrophilic because they expose a large number of polar amino acid residues lining up perpendicularly to the surface, also causing higher surface roughness. Acidic or basic treatments also have a great influence on the properties of zein-coated particles, as zein coatings have stronger binding affinity and higher mass adsorption on surfaces exposing carboxyl groups rather than alkyl groups (Wang et al., 2006). The assembly and structure of zein in solutions is a complex function of solvent polarity and zein charge.

Later, another study investigated changes in zein conformation as a function of solvent chemistry using CD analysis. The secondary and tertiary structure of zein protein could be varied by the type of solvent and the temperature (Selling et al., 2007). The secondary structure was weakly affected by ethanol to water ratio variation, while the tertiary structure slightly changed in the presence of acetic acid-water solutions. Reversible changes of the primary and tertiary structures were observed by heating zein solution in 80% v/v ethanol-water up to 70°C followed by cooling. With the aim to understand protein-solvent interaction and whether zein behaves as a polyelectrolyte or differently, SAXS, rheology and CD analyses were carried out on zein solutions in acetic acid-water (Li et al., 2011). Zein solutions in acetic acid behaved similarly to polyelectrolyte solutions in good solvents, as proved by molecular measures of protein folding, such as the concentration scaling behavior which was near the theoretical predictions for polyelectrolytes. Indeed, the folding or unfolding behavior of zein may change the surface of interaction with the solvents, leading to different dissolution. Afterwards, the same research group progressed their study on zein solution properties also in ethanol-water solvents (Li et al., 2012). Protonation of zein was identified as a key parameter for its solubilization. Hence, acetic acid-water was found to improve zein solubility respect to ethanol-water, because of the more favorable zein protonation in the presence of acetic acid. They also found out that zein molecule size increased by decreasing the content of ethanol in aqueous mixtures (from 80% to 70% v/v); the largest zein size was found in solutions containing acetic acid-water solvents suggesting that zein was more unfolded or swollen in the presence of acetic acid. As predicted by the polyelectrolyte behavior, with increasing zein concentration upon 43 mg/mL, zein size slightly decreased in solutions containing acetic acid-water: this behavior was due to overlapping and mutually penetrating behavior, because of crowding and repulsive long-range electrostatic interactions among zein chains. Another study on zein conformational changes in solution was carried out more recently (Erickson et al., 2020). CD spectra of zein solutions showed that, by increasing water in ethanol-water solvents, random coils and β -sheets began to appear instead of α -helix structures. This result was confirmed by molecular dynamics simulation performed on zein solutions in 45% and 75% v/v ethanol-water for the validation of two α -zein molecular models: the hairpin-like (Figure 1C) and the triple-helix models (Figure 1D). The hairpin-like model best represented zein conformation with increased β -sheets in 45% v/v ethanol-water.

TABLE 2 List of aqueous binary solvents (Lawton, 2002) and their greenness level according to Pfizer guide (Byrne et al., 2016; Joshi and Adhikari, 2019).

| The solvent in combination with water | Greenness level |
|---------------------------------------|-----------------|
| Acetone | Usable |
| Acetonyl acetone | Usable |
| n-Butanol | Usable |
| t-Butanol | Usable |
| s-Butanol | Usable |
| Dioxalane | Usable |
| Dioxane | Undesirable |
| Ethanol | Usable |
| Isobutanol | Usable |
| Isopropanol | Usable |
| Methanol | Usable |
| n-Propanol | Usable |

SEM analysis showed spherical zein aggregates (0.1–10 μm) in zein solution with 45% v/v ethanol-water and this could be explained by the increase in β -sheets occurring at higher water content.

A recent investigation on zein solubility in ethanol-water solvents evidenced the influence of amino acid composition of zein fractions (α -, β -, γ -, and δ -zein) on the physicochemical properties and exposed functionalities of zein based products prepared from those solutions (Gao et al., 2021). In detail, the three zein fractions allowed the successful production of NPs from ethanol-water solvents, with different characteristics in terms of wettability, hydrodynamic size and surface hydrophobicity. β -zein based NPs showed poor surface lipophilicity, as measured through an optical contact angle meter, using medium-chain triglyceride as the oil phase. α -zein NPs showed the highest surface hydrophobicity, leading to high oil-water interfacial adsorption speed. γ -zein based NPs showed low surface lipophilicity and the highest hydrodynamic size (383 \pm 26 nm) compared to α -zein (207 \pm 12 nm) and β -zein NPs (155 \pm 9 nm). Furthermore, respect to α -zein and β -zein, γ -zein contains more cysteine residues forming a crosslinked structure by disulfide bonds.

Recently attention is focused on the use of the most environmentally friendly and sustainable solvents for zein. As water represents the greenest solvent, aqueous binary solvents represent the best choice for zein solubilization under environmentally-friendly conditions. Table 2 reports a list of aqueous binary mixtures used for zein solubilization (Lawton, 2002) together with their level of greenness according to the Pfizer guide (Alfonsi et al., 2008; Byrne et al., 2016). Other solvent selection guides have also been developed to classify solvents based on environmental impact, safety, waste management, effects on human health or life cycle, in order to state which solvents can be considered 'greener' and which ones should be replaced because of significant issues (GSK, 2010; ACS-GCI, 2011; Henderson et al., 2011; Prat et al., 2013). A comparison among the different guides shows that they are in agreement, with

only slight discrepancies in few cases. Interestingly, all solvents used in aqueous binary mixtures for zein solubilization are classified as “usable”, except for dioxane.

Considering the wide range of possible solvents available for zein solubilization, a thorough understanding of zein molecule assembly and their phase behavior in these solvents is still lacking.

3.2 Methods for the preparation of zein nanoparticles

Various methods have been employed for the preparation of zein NPs for drug delivery. The selection of the method depends on solvent types and ratio, which are crucial to obtain specific characteristics for the nanocarriers.

Zein NPs have been generally prepared from binary solvent systems containing water, using a method that exploits the solubility of zein in various solvents and that has been firstly called *phase separation method*, but investigations in literature report also other names such *anti-solvent precipitation* (Cheng et al., 2019), *liquid-liquid dispersion* (Zou et al., 2012) and *coacervation* (Madan, 1978; Arshady, 1990; Swarbrick, 2007) methods (Yan et al., 2022). This technique induces NP formation by the dispersion of a primary phase, where zein is soluble, into a non-solvent for zein (typically water) which is miscible with the solvent. It is characterized by basic experimental settings and steps, is low-cost and usually shows high drug encapsulation efficiency, so represents the most widespread method to prepare zein NPs (Pascoli et al., 2018). *Liquid-liquid dispersion* has been first reported for the preparation of micro-rods (Alargova et al., 2004; Alargova et al., 2006): rods were obtained by mixing a polymer solution into an organic solvent, which was unable to dissolve the polymer while was miscible with the solvent. The same principle was then applied for the production of zein NPs (Zhong and Jin, 2009). According to this method, zein dissolved in an aqueous binary mixture (organic phase) was added to an anti-solvent (aqueous phase) under stirring. Due to the good miscibility of the used organic solvent with water, the inter-diffusion mechanism of solvents reduces the solubility of zein, leading to its precipitation into particles. Process parameters such as solvent ratio, method of injection (drop-wise or continuous addition), and stirring speed and method (magnetic stirring, vortex, sonicator, etc.) highly affect the results, such as NP size and morphology, leading to low experimental reproducibility and scalability for large volumes. Zhong and Jin conducted an interesting research on the effects of different process parameters (shear stress, aqueous ethanol ratio and zein concentration) on zein NPs (Zhong and Jin, 2009). Effects of shear stress, by applying different stirring speeds (5,000, 10,000 and 15,000 rpm) were studied, while keeping constant ethanol to water ratio and zein concentration. DLS and SEM showed that the use of lower speeds allowed NP dispersion into droplets with spherical shape and higher sizes (smaller than 200 nm). The influence of ethanol concentration increase from 55% to 90% on NP features was evaluated at the fixed stirring speed of 5,000 rpm. DLS and SEM analyses were then performed, showing that higher ethanol concentrations led to lower NP diameters. This result was due to NP formation kinetics: higher initial ethanol concentration in the solvent was associated with a slower decrease

of ethanol concentration in the dispersed droplets till the solubility limit could be reached. During this time, zein droplets could be sheared and broken-up to achieve smaller sizes before precipitation and solidification. Finally, the impact of zein concentration from 6.5 to 300 mg/mL was evaluated. DLS analysis showed that lower zein concentrations allowed the preparation of smaller NPs; on the other hand, at higher concentration than 100 mg/mL large agglomerates formed. This behavior depends on the fact that by decreasing zein concentration, zein solution viscosity also decreased and zein droplets could be more easily deformed, allowing the formation of smaller NPs.

Recently, innovative *dialysis methods* have been studied as alternatives to traditional approaches. Rodríguez-Félix et al. developed an antisolvent-dialysis method in order to form zein micro- and nanoparticles through an easy, one-step and reproducible process (Rodríguez-Félix et al., 2020). Despite being innovative, this method is also highly eco-friendly in terms of low energy consumption and waste production, due to the simultaneous combination of zein purification from hydrophobic xanthophylls, the yellow pigments of zein. This leads to the production of white zein particles without the need of a second processing step, thus decreasing the amount of solvent needed for both operations. This allows a reduction in the amount of solvents required for carrying out the operations separately. Figure 2 shows a schematic representation of the antisolvent-dialysis method. Basically, zein solution is put into a dialysis membrane inside a dialyzer, filled with distilled water at 1:10 ratio under magnetic stirring for 12 h at room temperature. This method induces zein self-assembly by the concentration gradient and polarity change of the solvents: the diffusion kinetics of the organic solvent through the dialysis membrane slow down gradually up to the nucleation of zein, with the formation of micro- and nanoparticles. This method allowed the production of a mixed population of microparticles ($0.74 \pm 0.2 \mu\text{m}$) and NPs ($243.2 \pm 94.5 \text{ nm}$). The authors proposed to apply these particles as natural and edible dyes for coatings in substitution to inorganic dyes or as polymeric matrices protecting molecules of interest in food industry.

Chen et al. performed a *multi-frequency ultrasound-assisted dialysis* to produce curcumin-loaded zein-sodium caseinate NPs (Chen et al., 2022b). They dissolved 1% w/v zein and sodium caseinate in 1,2-propanediol aqueous solution at 80% v/v by stirring and centrifugation to remove larger particles. Then zein-sodium caseinate solution was put into a dialysis membrane inside a dialyzer filled with deionized water in a 1:2 ratio, and the entire system was placed into an ultrasonic bath reactor (20–28–40 Hz, 300 W, 25°C, 10 min). After this process, dialysis was completed at room temperature to obtain the effective formation of stable zein NPs. Results showed that the optimal frequency of 20/40 kHz allowed the production of uniform and regular spherical NPs with $225 \pm 4 \text{ nm}$ size, low polydispersity index and negative zeta potential. The sonication process together with the use of sodium caseinate improved storage and thermal stability of NPs, and allowed high encapsulation efficiency and antioxidant activity of curcumin. They finally demonstrated controlled curcumin release under simulated gastrointestinal digestion conditions, confirming the success of the multi-frequency ultrasound-assisted dialysis technique and the possibility to extend its application to other hydrophobic compounds.

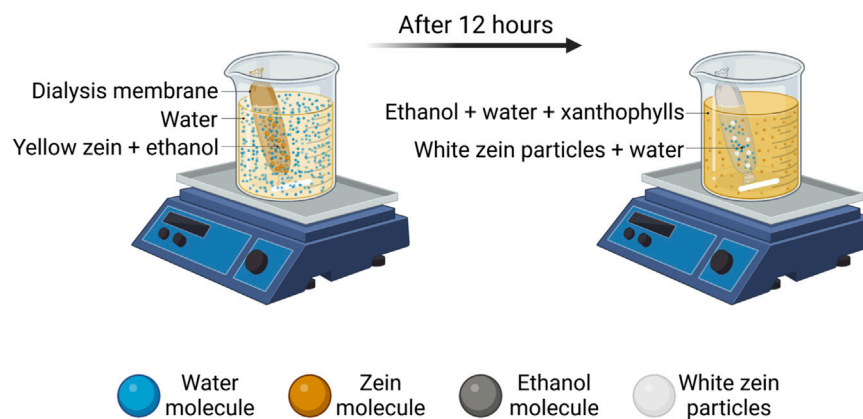


FIGURE 2

Antisolvent-dialysis method for the production of white zein micro- and nanoparticles (Rodríguez-Félix et al., 2020). Created with BioRender.com.

Few studies focused on the production of zein NPs through unconventional scalable *electrospraying* and *supercritical antisolvent precipitation techniques*.

Electrospraying is a single-step method that is based on the use of an electric field to induce the separation of the zein solution (extruded through a metal nozzle) into charged droplets. The principle is the same as for the electrospinning technique, but instead of obtaining nanofibers, it is possible to produce monodispersed NPs with different characteristics by modulating the electric voltage, solution concentration and flow rate, and nozzle-collector distance. Gomez-Estaca et al. obtained curcumin-loaded zein NPs with sizes ranging from 175 to 900 nm increasing polymer concentration (Gomez-Estaca et al., 2012). They found unaltered morphology of NPs after curcumin encapsulation and the process also allowed high encapsulation efficiency (85%–90%). An implementation of this method is coaxial electrospinning, which allows to produce hybrid NPs (Kang et al., 2021). Kang et al. prepared drug loaded polymer-lipid hybrid NPs loaded with tamoxifen citrate (a hydrophobic anticancer drug) by electrospinning. The drug-zein solution in aqueous ethanol at 70% v/v was exploited as the core working fluid while a solution of lipid in pure ethanol was exploited as the shell fluid. Zein core-lipid shell nanostructures with a round shape were obtained. The major disadvantages of this technique are the high cost and energy consumption which limit its application in the pharmaceutical, biotechnology, and medical industries as competitive drug delivery systems.

Supercritical antisolvent precipitation techniques (SAS) consists of a feed, continuously sprayed into supercritical CO₂ that acts as an anti-solvent for the majority of polymers (Franco and Marco, 2020). Rosa et al. studied the feasibility of SAS for the co-precipitation of a vitamin complex containing riboflavin, δ -tocopherol and β -carotene in zein microcapsules (Rosa et al., 2020). They obtained spherical microcapsules with a size varying in the 8–18 μ m range depending on which vitamin was encapsulated. Despite the scalability of the process, SAS is incompatible with several pharmaceuticals and requires a specific and expensive setting. Therefore *phase separation* conventional method remains the most applied for the preparation of zein NPs.

3.3 Zein nanoparticles for drug delivery

Protein-based NPs usually show biocompatibility, biodegradability and minimization of hypersensitivity reactions (Mishra et al., 2010). Furthermore, protein-based NPs are generally prepared avoiding the use of toxic chemicals or organic solvents (Lohcharoenkal et al., 2014). Zein protein NPs have been mostly exploited for the encapsulation of hydrophobic compounds (e.g., essential oils); their large amount of nonpolar amino acid residues favor zein interaction with hydrophobic drugs to achieve high drug loading efficiency. However, in a few cases, hydrophilic drugs, including oligonucleotides, have also been encapsulated into zein NPs. The majority of research studies prepared zein NPs using zein solutions in binary solvents containing water which is the greenest solvent.

The following paragraphs describe the preparation of zein NPs loaded with hydrophobic and hydrophilic drugs using zein solutions in different solvent mixtures. The dependence of NP size and physicochemical properties on used binary aqueous solvents and ratio between them is also highlighted. Table 3 collects main features of exemplary zein NPs discussed in this paper.

3.3.1 Hydrophobic drug delivery by zein nanoparticles

Over the years, zein has been mainly studied as a nanocarrier for the release of hydrophobic compounds and NPs have been generally prepared from solutions in ethanol-water solvents (Tran et al., 2019) (Table 3). One advantage of zein NPs over other carriers is their superior stability in the gastric and intestinal tract upon oral administration thanks to its isoelectric point that allows to maintain stability against aggregation of zein NPs at acidic pH. Furthermore, zein shows resistance to degradation at acidic pH and in the presence of proteolytic enzymes. These characteristics allow the use of zein NPs as carriers for drugs that need to be protected during oral delivery against the harsh gastrointestinal conditions (Lee et al., 2013; Cheng et al., 2019).

Therefore, zein NPs are often exploited for oral delivery of compounds to prevent their degradation in the harsh gastrointestinal tract (GIT) conditions because at acidic pH they remain stable against aggregation.

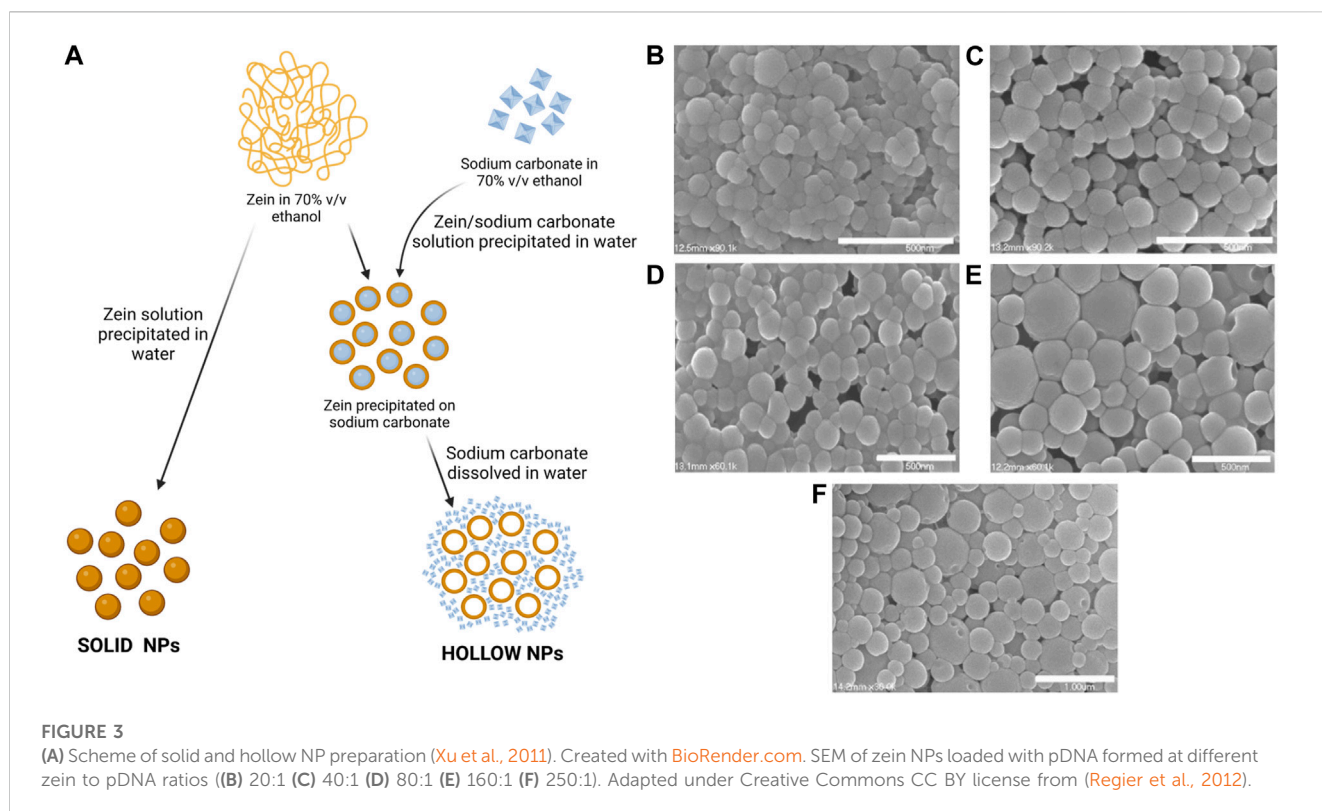
TABLE 3 Main information on zein NPs discussed in this review paper.

| Solvent | Method | Encapsulated compound | Size [nm] | Application | Reference |
|--------------------|----------------------------|-----------------------------------|--|---|--|
| Dimethyl sulfoxide | Phase separation method | Maytansine | 112 | Lung cancer | Yu et al. (2020) |
| 90% ethanol | Coacervation | PTEN and TRAIL genes | 133–239 | Hepatocellular carcinoma | El Sharkawi et al. (2017) |
| 85% ethanol | Phase separation method | Oregano, red thyme, and cassia | 50–125 | Antioxidant, antimicrobial | Parris et al. (2008) |
| 80% ethanol | Antisolvent-dialysis | - | 243–740 | Polymer matrix to protect molecules of interest in food, natural colorant | Rodríguez-Félix et al. (2020) |
| | Anti-solvent precipitation | Lutein | 75 | Heart disease, stroke and cataract formation | Cheng et al. (2019) |
| | Liquid-liquid dispersion | Cranberry procyanidins | 392–447 | Urinary infections | Zou et al. (2012) |
| | Phase separation method | Catalase and superoxide dismutase | 255 | Rheumatoid arthritis | Lee et al. (2013) |
| | Nanoprecipitation method | Chloroquine | 172–186 | Apinal cord injury | Chen et al. (2022c) |
| 70% ethanol | Phase separation method | 5-fluorouracil | 115 | Liver | Lai and Guo (2011) |
| | Phase separation method | Metformin | 63–102 for hollow NPs, 135–160 for solid NPs | Anti-diabetic | Xu et al. (2011) |
| | Phase separation method | Doxorubicin | 200–240 | Anti-cancer | Dong et al. (2016) |
| | Liquid-liquid dispersion | Thymol and carvacrol | 330 | Antioxidant, antimicrobial | Wu et al. (2012) |
| | Phase separation method | Prednisolone | 374–15117 | Bowel disease | Lau et al. (2012) |
| | Phase separation method | Hydrocortisone, mesalazine | 442–1,100 for hydrocortisone, 1,381–12730 for mesalazine | Bowel disease | Lau et al. (2013) |
| | Coacervation | pDNA | 158–397 | Kidney | Regier et al. (2012) |
| 55%–90% ethanol | Liquid-liquid dispersion | - | 100–200 | Drug delivery | Zhong and Jin (2009) |
| 90% isopropanol | Liquid-liquid dispersion | Fish oil | 350–450 | Antioxidant | Zhong et al. (2009) |
| 80% acetone | Emulsion-diffusion method | - | 135 | Botany | Prasad et al. (2017), Ristroph et al. (2017) |
| | Emulsion-diffusion method | - | 145 | Botany | Bonser et al. (2020) |
| | Emulsion-diffusion method | Methoxyfenozide | 209 | Botany | Hanna et al. (2022) |

Cheng et al. prepared zein NPs encapsulating lutein, a carotenoid compound (a xanthophyll) naturally present in corn and also in leafy green vegetables, and known to have beneficial effects on reducing the risk of heart disease, stroke and cataract formation (Cheng et al., 2019). However, bio-accessibility of lutein naturally present in zein and other plant derivatives is low (14%–55%) (Serrano et al., 2005; Reboul et al., 2006). Zein NPs represent valuable carriers for lutein delivery, protecting this compound from heat, oxygen and light. In this work, lutein-loaded zein NPs with an average size of 75 nm were prepared by anti-solvent precipitation, dissolving zein (25 mg/mL) in 80% v/v aqueous

ethanol solution and lutein (0.5 mg/mL) in ethanol. NPs were formed by drop-wise dispersion in water.

Zou et al. produced zein NPs encapsulating cranberry procyanidins (CPs) through a liquid-liquid process (Zou et al., 2012). Procyanidins have antioxidant properties, thus reducing the risk of cardiovascular diseases and cancers. Among them, cranberry procyanidins show beneficial effects against urinary infections (Foo et al., 2000; Santos-Buelga and Scalbert, 2000; Prior and Gu, 2005). Procyanidins have a high binding affinity to proteins rich in prolines and they are usually employed to remove excessive tannins from wines. Since zein has a proline content of



around 10%, innovative CPs-loaded zein NPs were prepared in order to improve the oral bioavailability of CPs. Different concentrations of CPs and zein (1 mg/mL) were dissolved in 80% v/v ethanol-water, then water was added, causing NP formation. By increasing CPs to zein weight ratio (from 1:8 to 1:2 wt/wt), the hydrodynamic diameter was found to increase from 392 nm to 447 nm, while zeta-potential decreased from 20.9 mV to 14.2 mV. The encapsulation efficiency increased by decreasing CPs to zein weight ratio, from 10% to 86%. CPs-loaded NPs showed superior *in vitro* biocompatibility compared to CPs alone, as assessed through *in vitro* cell tests with human promyelocytic leukemia HL-60 cells.

Zein NPs encapsulating 5-fluorouracil (5-FU) were also developed for intravenous delivery to target liver tissue (Lai and Guo, 2011). Zein was selected to protect 5-FU from enzymatic degradation and to shield its potential toxicity. NPs were prepared by the phase separation method. 5-FU and zein (12.5 mg/mL) were dissolved in 70% v/v ethanol-water, then added to distilled water. The hydrodynamic diameter was 114.9 ± 59.4 nm, zeta potential was -45 ± 0.3 mV and encapsulation efficiency was $60.7\% \pm 1.74\%$. Initial burst release was due to weakly bonded drug and attributed to zein NP core-shell structure. Indeed, during the synthesis process zein first forms a core, and then slowly adsorbs free zein to form a shell under a layer-by-layer assembly architecture. Therefore, burst release of drug may be probably due to the release of the drug portion present in the shell or at the core-shell interface. *In vitro* release was complete within 24 h.

Xu et al. prepared hollow zein NPs for various drug delivery applications (Figure 3A) (Xu et al., 2011). The anti-diabetic drug metformin was used as a drug model. Hollow NPs were produced by phase separation method dissolving zein (91 mg/mL) and sodium

carbonate crystal (7 mg/mL) in 70% v/v ethanol-water. Then, this solution was precipitated into water using different volumes. Since sodium carbonate is highly soluble in water, the sacrificial cores immediately dissolved after zein precipitation on its surface, forming hollow NPs. After, an aqueous solution of metformin was added to the NP dispersion for 5 h at 50°C. Control solid zein NPs were prepared by precipitating zein solution in ethanol-water (12 mg/mL) into water. The size of both hollow and solid NPs presented a decreasing trend (from 102.3 nm to 62.9 nm for hollow NPs and from 160 nm to 135 nm for solid NPs) by increasing water to zein solution volume ratio. Hollow NPs showed lower sizes than solid NPs probably due to the presence of sodium carbonate nanocrystals acting as heterogeneous nucleation sites, and limiting the increase of diameters. Hollow NPs showed slower release kinetics and released a higher overall drug amount respect to solid NPs. *In vitro* biocompatibility and cell internalization was assessed using NIH 3T3 mouse fibroblast cells.

Yu et al. described maytansine (DM1)-loaded zein NPs prepared by phase separation method using dimethyl sulfoxide (DMSO) solvent (Yu et al., 2020). DM1 has high and broad antitumor activity. DM1 and zein (60 mg/mL) were dissolved separately in a DMSO solution, then mixed together and dropped into distilled water while stirring. NPs showed a hydrodynamic diameter of 112 nm, a polydispersity index of 0.21, a zeta potential of 37 mV and high encapsulation efficiency (83%). *In vitro* release kinetics was characterized by a burst release within the first 8 h of about 20%, followed by a release of 40% after 24 h (Li et al., 2017; Sabra et al., 2018). DM1-loaded zein NPs were stable in Fetal Bovine Serum (FBS) *in vitro*. High cellular uptake of NPs was demonstrated with perinuclear disposition and strong inhibitory effect on A549 cells.

Successful lung tumor treatment and nontoxic effects were also demonstrated by *in vivo* mouse tests.

3.3.2 Essential oils delivery

Zein NPs have been widely investigated for the loading and release of essential oils. Essential oils are hydrophobic and volatile compounds derived from plants, with a broad spectrum of biological activities, including antifungal, insecticidal, antimicrobial, and antioxidant properties. Examples include oregano, thyme, lavender and lemon oils.

To the best of our knowledge, the first work on essential oils encapsulation into zein NPs was published by Parris et al. (Parris et al., 2008). Three essential oils (oregano, red thyme, and cassia) were loaded into zein NPs by phase separation method. Zein (66.7 mg/mL) and essential oils were dissolved in an aqueous ethanol solution (85% v/v), then dispersed in water. Zein NPs encapsulated 20% oregano and thyme, and 13% cassia oil.

Later, another study focused on the encapsulation of two essential oils (thymol and carvacrol) into zein NPs, using liquid-liquid dispersion method (Wu et al., 2012). Essential oils and zein (50 mg/mL) were dissolved in 70% v/v aqueous ethanol solution and then subjected to shear stresses in water through homogenization. NPs showed an average hydrodynamic size below 330 nm and a polydispersity index below 0.3. Encapsulation efficiency was more than 60%, with higher thymol loading compared to carvacrol although with no significant differences. The authors hypothesized that encapsulation efficiency, measured after lyophilization, was slightly lower than the real value, due to the volatility of essential oils. Although carvacrol has a higher solubility in water, no difference in the release behavior of the two tested essential oils was detected.

In another study, zein NPs were found to encapsulate more efficiently non-volatile fish oil (above 90%) than other volatile essential oils (Zhong et al., 2009). Fish oil was loaded into zein NPs via liquid-liquid dispersion (Shaw et al., 2007). In this case, 90% v/v isopropanol was used to dissolve zein (40–67 mg/mL) and fish oil. Even if isopropanol is not a GRAS solvent, it was used because fish oil has a reduced solubility in ethanol solutions. The isopropanol percentage was selected taking into account zein insolubility at higher isopropanol concentrations. NP sizes ranged between 350 and 450 nm, and did not show a round morphology differently from zein NPs prepared in ethanol-water solutions. Lyophilized NPs were able to protect loaded fish oil from oxidation.

3.3.3 Hydrophilic drug delivery by zein micro- and nanoparticles

Zein NPs encapsulating water-soluble compounds have been less investigated for oral drug delivery. Lau et al. extended the use of a coacervation method, previously developed for the preparation of zein microparticles loaded with prednisolone, a hydrophobic anti-inflammatory drug, to the encapsulation of hydrophilic agents (Lau et al., 2012). By this method, they prepared zein microparticles loaded with two drugs for the treatment of inflammatory bowel disease: a hydrophobic drug with similar chemical structure to prednisolone, called hydrocortisone, and a structurally different water-soluble drug, called mesalazine (Lau et al., 2013). Zein microparticles were prepared by dissolving zein into 70% v/v

aqueous ethanol solution. They also optimized the method for solution stirring, usually not well described in published protocols, with the aim to enhance microparticle reproducibility and drug encapsulation efficiency. In details, vortex mixer allowed higher drug encapsulation efficiency compared to magnetic stirring. Mesalazine-loaded zein microparticles showed higher loading efficiency but lower reproducibility compared to hydrocortisone-loaded zein microparticles.

Another study explored the encapsulation of two antioxidant proteins, catalase and superoxide dismutase (SOD), as scavengers for reactive oxygen species (ROS) generated by macrophages in rheumatoid arthritis (Lee et al., 2013). These two proteins were loaded into zein NPs for oral delivery to protect them from harsh environment in the gastrointestinal tract. Glisodin[®], the first effective oral delivery system for SOD release is based on gliadin, a prolamin from wheat, which may induce Celiac disease. Therefore, authors prepared NPs based on an alternative prolamin, zein, for the delivery of the same proteins. NPs were prepared by phase separation method by dissolving zein (55 mg/mL) in 80% v/v ethanol-water, then adding the therapeutic proteins. NPs showed an average hydrodynamic diameter of 255 nm at pH 7.0 and the encapsulation efficiency was 31% for SOD and 45% for catalase. Zein conjugated with folic acid by an amide linkage to lysine residues allowed targeted drug release to activated macrophages, demonstrated by *in vitro* cell tests.

Dong and coworkers prepared zein NPs loaded with doxorubicin (DOX), a water-soluble anti-cancer drug, in order to provide effective drug dose with no cytotoxic effects (Dong et al., 2016). Zein (10 mg/mL) and DOX were dissolved in 80% v/v ethanol-water and NPs were produced by phase separation method. The average hydrodynamic diameter of DOX-loaded zein NPs ranged from 200 to 240 nm, the polydispersity index was around 0.15–0.20, zeta potential was -50 mV and the encapsulation efficiency was about 90%. *In vitro* release tests displayed a pH-responsive trend. Three pH conditions were tested (5.0, 6.5 and 7.4) and the cumulative release of zein NPs reached 69%, 83% and 60% after 100 h, respectively, thus achieving higher values at more acidic pH. The anti-proliferative effect of DOX-loaded NPs on HeLa cells was studied. NPs entered the cytoplasm of HeLa cells *via* micropinocytosis and DOX was released in a controlled and sustained manner from zein NPs.

More recently, zein NPs were selected for chloroquine loading, an anti-malarian drug employed to decrease inflammation after spinal cord injury (Chen X. et al., 2022). Zein (6.5 mg/mL) was dissolved in 80% v/v ethanol-water, then water and chloroquine were added separately. Chloroquine-loaded zein NPs showed a hydrodynamic diameter ranging from 172 to 186 nm, a low polydispersity index, negative zeta potentials between -30 and -40 mV and a high encapsulation efficiency (61%–77%). An *in vitro* controlled delivery of 4 days was monitored. *In vitro* tests using microglial BV2 cells showed a reduction of pro-inflammatory factors after zein NP treatment. Microglia polarization towards the M2 phenotype was enhanced by zein NPs. Indeed, zein contributed to the attenuation of inflammation thanks to the beneficial effects of its degradation products. *In vivo* local administration on rats caused improved functional recovery, inhibited neuronal apoptosis and suppressed neuroinflammation.

3.3.4 Delivery of oligonucleotides by zein nanoparticles

Non-viral gene therapy has received widespread attention for its ability to introduce foreign exogenous DNAs or RNAs into cells affecting cell behavior (e.g., replacing genes or regulating their expression) (Shim et al., 2018). Zein NPs protect oligonucleotide cargo from degradation and allow their controlled and sustained release for efficient transfection. However, the use of zein NPs for oligonucleotide encapsulation has been poorly investigated to date.

To the best of our knowledge, Regier et al. were the first ones to develop zein NPs encapsulating plasmid DNA (pDNA) by the coacervation technique for kidney targeting (Regier et al., 2012). Zein is an amphiphilic protein with a high content of polar and protonable glutamine residues and an isoelectric point of 6.2–6.8, allowing the encapsulation of negatively charged pDNA *via* electrostatic interaction with positively charged zein chains, upon pH regulation. Zein (10 mg/mL) was dissolved in 70% v/v ethanol-water at pH 3, then pure ethanol at pH 3 was added, followed by addition of pDNA aqueous solution. Depending on zein to pDNA ratio, prepared NPs showed sizes ranging from 158 to 397 nm (Figures 3B–F) and zeta potential values ranging from –20 mV to –50 mV. Degradation tests in the presence of DNase I confirmed integrity of loaded pDNA. Zein NPs showed sustained but incomplete *in vitro* pDNA release in PBS within 7 days, with only 18% of loaded pDNA released, after an initial burst release within the first 12 h. *In vitro* cell tests showed biocompatibility, and efficient NP internalization by Caco-2 (Human colon carcinoma cells) and HEK 293 T (Human embryonic epithelial kidney cells) cells. Indeed, the N-terminal region of γ -zein, rich in proline, probably served as a peptide carrier interacting with the phosphate diester anionic polar heads of the cell membrane, favoring the endocytosis of zein NPs, with pDNA transfer into the cell cytoplasm (Fernández-Carneado et al., 2004). Interestingly, cell internalization could be analyzed by confocal microscopy exploiting zein autofluorescence (with excitation wavelengths at 365–395 nm and 465–485 nm).

A few years later, zein NPs were employed again in gene therapy, for the encapsulation of pDNAs encoding Phosphatase and Tensin homolog (PTEN) and Tumor Necrosis Factor - Related Apoptosis-Inducing Ligand (TRAIL) for the treatment of hepatocellular carcinoma (El Sharkawi et al., 2017). Zein NPs were prepared by the coacervation method with slight modifications compared to the previous protocol. Zein (20 mg/mL) was dissolved in highly concentrated ethanol solution (90% v/v) at pH 3 and then pDNA solution at pH 4 was added. The average NP hydrodynamic diameter and zeta potential ranged from 132 to 238 nm and from –37 to –24 mV, respectively. Encapsulation efficiency varied from 12% to 42%. *In vitro* tests showed decreased cell viability and proliferation rate of liver tumor cells (HepG2) treated with NPs. *In vivo* experiments in rat tumor model confirmed a significant decrease of liver metastasis by increased mRNA expression of p53 (a marker of apoptosis) and decreased mRNA levels of VEGF and MMP-2 (markers for angiogenesis and metastasis, respectively), measured from liver homogenates.

Overall, only a few studies have been published on the use of zein NPs delivering hydrophilic therapeutic compounds. Hence, further research is needed to optimize the preparation of such NPs with the aim to obtain higher encapsulation efficiency.

3.3.5 Greener preparation methods for zein nanoparticles

The herein performed literature survey on zein NPs has shown that phase separation method using ethanol-water (Table 3) is the most frequently used approach. As underlined at Par. 3.2, this is due to the simple experimental settings and low-cost of the technique. However, ethanol-water binary solvents are suboptimal considering their environmental impact.

Greener phase preparation methods should be developed, for example, by exploiting acetone-water solvents. Acetone has been defined as the least toxic industrial solvent (Joshi and Adhikari, 2019). However, only a few examples of zein nanocarriers prepared from acetone-water solutions have been published for applications in the botany field (Table 3) with the aim to address the challenge of controlled delivery of pesticides and fertilizers to plants. Indeed, in the field more than 90% of conventional agrochemicals are unable to reach the desired target using traditional administration methods (Pérez-Guzmán and Castro-Muñoz, 2020). In botany, zein NPs delivering pesticides and fertilizers have been proposed by exploiting their adhesion to the root system or the foliage (Prasad et al., 2017). Prasad et al. prepared positively charged zein NPs for pesticide delivery and evaluated their uptake and translocation to sugar cane plants through plant roots (Prasad et al., 2017). Zein NPs were prepared by dissolving zein (40 mg/mL) in 80% v/v acetone-water solution, followed by its dropping into an aqueous phase (containing a positively charged stabilizing surfactant for the specific purpose of this work). Spherical NPs with a hydrodynamic size of 135 nm and a polydispersity index of 0.20 were obtained (Table 3). Later, the same research group studied the biodistribution of zein NPs, from the root to the leaves of soybean plants, with the aim to widen the application of zein NPs for pesticide delivery to different crops (Ristroph et al., 2017). Zein NPs were prepared with slight variations respect to the previously developed protocol (32 mg/mL of zein dissolved in 80% v/v acetone-water) (Table 3). Further research by another group investigated the effects of positively-charged zein NPs, prepared by the same method described above, on three models of larval noctuid pests, as a potential alternative strategy for pest control (Bonser et al., 2020) (Table 3). More recently, zein NPs were prepared by the same protocol and then surface modified with a negatively charged coating for the encapsulation and controlled delivery of a pesticide (Hanna et al., 2022) (Table 3).

Despite zein NPs prepared through greener conditions have been applied in botany only, their use could be extended to other therapeutic applications in regenerative medicine.

3.3.6 Zein-based core-shell nanoparticles

Although zein has been frequently used alone for the preparation of drug-loaded micro- and nanoparticles, a great number of studies have also focused on the preparation of zein-based core-shell NPs in combination with another biomaterial (Demir et al., 2017). Two applications have been investigated: 1) the use of zein as coating on NPs based on another material, exploiting zein favorable stability at acid pH and ability for sustained release; 2) the use of zein as core for NPs coated with a different material, in order to overcome colloidal instability of zein NPs (Yuan et al., 2022b).

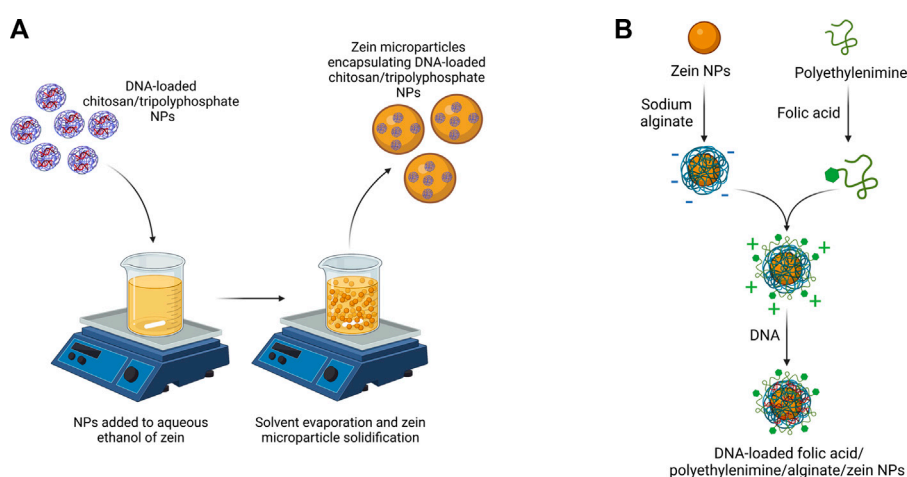


FIGURE 4

(A) Scheme of DNA-loaded chitosan/tripolyphosphate NPs coated with zein (Farris et al., 2017) (B) Scheme of the preparation of DNA-loaded zein NPs coated with alginate, poly (ethyleneimine) and folic acid (Chen Y. et al., 2022). Created with BioRender.com.

As a relevant example of zein-coated NPs, Luo et al. prepared chitosan/tripolyphosphate NPs loaded with selenite and coated with zein (Luo et al., 2010). Zein coated NPs maintained nanoscale dimensions (200–400 nm), showed improved encapsulation efficiency (with an increase from 60% to 95% after zein coating) and release profile (with reduced burst release within the first 4 h in PBS). A few years later, chitosan/tripolyphosphate NPs were employed again for oral gene delivery and coated with zein (Farris et al., 2017). As shown in Figure 4A, DNA-loaded chitosan/tripolyphosphate NPs were encapsulated in zein microparticles. Zein coating caused an increase in DNA encapsulation efficiency and a slower *in vitro* release kinetics in simulated gastric fluid compared to DNA-loaded chitosan/tripolyphosphate NPs. *In vivo* tests in mouse model confirmed successful cell transfection. More recently, the introduction of synthetic PLGA NPs coated with zein was proposed for the release of docosahexaenoic acid (Liu et al., 2021).

A recent review paper has discussed the effect of composition, preparation and storage methods of zein-based NPs with zein core on their stability (Yuan et al., 2022a). Different materials have been proposed as coatings for zein NPs, such as polysaccharides, proteins and surfactants (Li and Yu, 2020). Such NPs have been prepared by simple methods, such as anti-solvent precipitation, layer-by-layer coating or by exploiting zein properties, such as surface charge tunability with pH. In some cases, polysaccharide-coated zein NPs were not stable, due to steric hindrance between polysaccharide coatings and functional moieties on zein NPs. This issue was not detected with alginate coating which has been deeply investigated in literature as stabilizer for NPs. For example, alginate has been used both alone for doxorubicin delivery (Ye et al., 2022a), or in combination with other charged polymers to generate stable complexes for DNA and folic acid (Figure 4B) (Chen Y. et al., 2022) or curcumin delivery (Yao et al., 2018; Liu et al., 2019). Propylene glycol alginate, an alginate derivative, was combined with zein to form a noncovalent binary complex for the release of quercetagenin (Sun et al., 2016), β -carotene (Wei et al., 2018) and

resveratrol (Wei et al., 2019). Other examples include soybean shell/zein NPs for quercetin release, exploiting soybean coating stability to pH (in the two to eight range), ionic strength and temperature (Li H. et al., 2019). Same NPs were applied for lutein release (Li et al., 2020). Zein-carrageenan core-shell NPs were successfully loaded with curcumin and piperine through a combination of electrostatic attraction, hydrogen bonding, and hydrophobic interactions (Chen et al., 2020). Pectin coating was also exploited to avoid agglomeration of zein NPs, loaded with tannic acid (Liang et al., 2021). In another application, ethanol-soluble pectin from sugar beet was exploited as a coating stabilizing zein NPs and favoring the delivery of curcumin (Zhang et al., 2022). Literature also reports coatings with pectin together with sodium caseinate (Chang et al., 2017a; Chang et al., 2017b). Hyaluronic acid was also applied as coating of zein NPs for quercetagenin (Chen et al., 2019b) and tetrahydrocurcumin (Zhu et al., 2022) delivery. Chitosan coating was applied on zein based NPs for the release of *a*-tocopherol (Luo et al., 2011), retinol (Park et al., 2015), quercetin (Ma et al., 2018), curcumin (Baspinar et al., 2018; Li et al., 2018) and folic and caffeic acid (Wusigale et al., 2021). Alternatively, carboxymethyl chitosan coating was used for zein NPs encapsulating vitamin D3 (Luo et al., 2012) and natamycin (Lin et al., 2020). Blend coatings applied on zein NPs include: i) chitosan/alginate for resveratrol delivery (Khan et al., 2019); ii) chitosan/dextran sulfate for the release of crocin carotenoid for Alzheimer's disease treatment (Song et al., 2022); iii) chitosan/hyaluronic acid for the co-delivery of curcumin and piperine (Chen et al., 2019a); iv) chitosan hydrochloride/sodium caseinate for thymol delivery (Zhang et al., 2014). Indeed, sodium caseinate is the mostly studied zein electrosteric stabilizer: it is an amphiphilic stabilizer which is adsorbed on the surface of zein NPs to provide an electrostatic and steric repulsion against aggregation (Patel et al., 2010; Luo et al., 2013; Ahmed et al., 2015; Pan and Zhong, 2016; Li et al., 2017; Zhang and Han, 2018; Chen et al., 2022a).

Concerning the use of surfactants as coatings for zein NPs, an interesting study compared zein based NPs and PLGA NPs for their

TABLE 4 Main literature examples of core-shell NPs with zein used as core or shell.

| Zein role in NPs | Composition of NPs | Encapsulated compound | Reference |
|---------------------------|--|------------------------|------------------------|
| Shell | Zein/chitosan/tripolyphosphate NPs | Selenite | Luo et al. (2010) |
| | Zein/chitosan/tripolyphosphate NPs | pDNA | Farris et al. (2017) |
| | Zein/PLGA NPs | Socosaheaxaenoic acid | Liu et al. (2021) |
| Core | Alginate/zein NPs | Doxorubicin | Ye et al. (2022a) |
| | Folic acid/polyethylenimine/alginate/zein NPs | pDNA | Chen et al. (2022d) |
| | Gelatin/alginate/zein NPs | Curcumin | Yao et al. (2018) |
| | Alginate/sodium caseinate/zein NPs | Curcumin | Liu et al. (2019) |
| | Propylene glycol alginate/zein NPs | Quercetagetin | Sun et al. (2016) |
| | Propylene glycol alginate/zein NPs | β -carotene | Wei et al. (2018) |
| | Propylene glycol alginate/zein NPs | Resveratrol | Wei et al. (2019) |
| | Soybean/zein NPs | Quercetin | Li et al. (2019b) |
| | Soybean/zein NPs | Lutein | Li et al. (2020) |
| | Carrageenan/zein NPs | Curcumin and piperine | Chen et al. (2020) |
| | Pectin/zein NPs | Tannic acid | Liang et al. (2021) |
| | Ethanol-soluble polysaccharide/zein NPs | Curcumin | Zhang et al. (2022) |
| | Pectin/sodium caseinate/zein NPs | - | Chang et al. (2017b) |
| | Pectin/sodium caseinate/zein NPs | Curcumin | Chang et al. (2017a) |
| | Hyaluronic acid/zein NPs | Quercetagetin | Chen et al. (2019b) |
| | Hyaluronic acid/zein NPs | Tetrahydrocurcumin | Zhu et al. (2022) |
| | Chitosan/zein NPs | α -tocopherol | Luo et al. (2011) |
| | Chitosan/zein NPs | Retinol | Park et al. (2015) |
| | Chitosan/zein NPs | Quercetin | Ma et al. (2018) |
| | Chitosan/zein NPs | Curcumin | Li et al. (2018) |
| | Chitosan/zein NPs | Curcumin | Baspinar et al. (2018) |
| | Chitosan/zein NPs | Folic and caffeic acid | Wusigale et al. (2021) |
| | Carboxymethyl chitosan/zein NPs | Vitamin D3 | Luo et al. (2012) |
| | Carboxymethyl chitosan/zein NPs | Natamycin | Lin et al. (2020) |
| | Chitosan/alginate/Zein NPs | Resveratrol | Khan et al. (2019) |
| | Chitosan/dextran sulfate/zein NPs | Crocin | Song et al. (2022) |
| | Chitosan/hyaluronic acid/zein NPs | Curcumin and piperine | Chen et al. (2019a) |
| | Chitosan hydrochloride/sodium caseinate/zein NPs | Thymol | Zhang et al. (2014) |
| | Sodium caseinate/zein NPs | - | Patel et al. (2010) |
| | Sodium caseinate/zein NPs | - | Luo et al. (2013) |
| | Sodium caseinate/zein NPs | Simvastatin | Ahmed et al. (2015) |
| | Sodium caseinate/zein NPs | - | Pan and Zhong (2016) |
| | Sodium caseinate/zein NPs | Rutin | Zhang and Han (2018) |
| Sodium caseinate/zein NPs | Curcumin | Chen et al. (2022a) | |
| Sodium caseinate/zein NPs | - | Li et al. (2017) | |

(Continued on following page)

TABLE 4 (Continued) Main literature examples of core-shell NPs with zein used as core or shell.

| Zein role in NPs | Composition of NPs | Encapsulated compound | Reference |
|------------------|---|-----------------------|-------------------------|
| | Sodium deoxycholate monohydrate or Poloxamer 188/zein NPs | Rutin | Gagliardi et al. (2021) |
| | Lecithin/zein NPs | - | Dai et al. (2016) |
| | Poloxamer 407/zein NPs | Dioscin | Zhang et al. (2021) |
| | Lysine/zein NPs | Glibenclamide | Lucio et al. (2017) |

ability to release rutin, with sodium deoxycholate monohydrate or poloxamer 188 as stabilizers (Gagliardi et al., 2021). The aim of the investigation was to compare physico-chemical characteristics of NPs prepared with the synthetic (PLGA) and the natural (zein) polymers, the capacity of PLGA or zein NPs to encapsulate and release rutin and NP antioxidant effect. Zein NPs were prepared by dissolving zein in 65% ethanol-water and adding sodium deoxycholate monohydrate or poloxamer 188. Results showed the enhanced ability of zein NPs to encapsulate rutin and release it in a controlled and sustained way, also thanks to the surfactant-based coating. Moreover, zein-based NPs showed increased *in vitro* antioxidant activity thanks to the synergistic effect of rutin and zein degradation products, compared to PLGA NPs. Other studies made use of other surfactants, such as lecithin (Dai et al., 2016) and poloxamer 407 (Zhang et al., 2021) for zein NP coating, while Lucio et al. prepared zein NPs in the presence of lysine as stabilizer (Lucio et al., 2017).

Table 4 collects the main hybrid core-shell zein NPs discussed in this paragraph.

3.3.7 *In vivo* immune response of zein nanoparticles

Beside safety, *in vivo* immunogenicity is a key parameter for *in vivo* application of particles, and it depends on their physicochemical properties and administration route. Zein is biocompatible and biodegradable. However, being a plant-derived protein, potential immune response *in vivo* may represent an issue. To date, studies on *in vivo* immune response induced by zein are still limited and have shown controversial results (Zhang et al., 2016). For example, zein microspheres were found to be immunogenic when administered intramuscularly and orally. Intramuscular administration of blank zein microspheres enhanced antibody reaction with immunoglobulin G (IgG), while oral administration increased immunoglobulin A (IgA) expression in the intestine area, keeping low IgG levels (Hurtado-López and Murdan, 2010). Some years later, Li and coauthors confirmed that zein NPs administered *via* intramuscular, subcutaneous and interpleural routes provoked enhancement of IgG antibodies in mice (Li F. et al., 2019). On the other side, non-immunogenic zein NPs (with hydrodynamic diameters ranging from 100 to 400 nm) were also patented (Perumal et al., 2009). The same authors showed that zein NPs with sizes above 400 nm, administrated *via* subcutaneous injection, caused an *in vivo* immune response in mice.

Very recently, another study for the repair of sciatic nerve defects in rat model made use of zein microparticles confirming that size is a discriminating parameter for immunogenicity. Authors of this study highlighted other influencing factors on

immunogenicity, such as the type of encapsulated drug (in their case, dexamethasone, an anti-inflammatory drug with immunosuppressive effects) and the porosity of the implanted structure, as it may inhibit neutrophil recruitment and enhance macrophage polarization towards the M2 phenotype. They studied the immunogenicity of zein compared to other materials, such as alginate, poly (lactic-co-glycolic acid) (PLGA) and polystyrene in rat model. They found out that zein first triggered an inflammatory reaction, similarly to non-degradable polystyrene; then, zein degradation occurred and the level of antigenicity was similar to that induced by alginate and PLGA, demonstrating zein safety (Yu et al., 2022).

Zein may be combined with polyethylene glycol (PEG) to inhibit immunogenicity. PEG is hydrophilic, highly water-soluble, anti-fouling and non-toxic. Moreover, it may improve zein colloidal stability in physiological environment. Two different methods have been investigated to produce zein-PEG NPs: PEG coating after NP formation and PEG chemical conjugation to zein protein before NP formation. PEG coatings can be easily obtained on zein NPs leading to improved serum stability, hemocompatibility, and good physical stability (El-Lakany et al., 2018; Reboredo et al., 2021; Ye et al., 2022b; Reboredo et al., 2022; Wiggers et al., 2022). However, PEG surface content is generally low and not homogeneous, hence it does not completely solve the issue of zein NP aggregation. On the other hand, zein chemical modification with PEG through carbodiimide reaction is laborious and requires longer time (zein functionalization needs days for reaction and dialysis). However, obtained NPs are advantageous for their low immunogenicity and improved serum stability (Podaralla et al., 2012; Soe et al., 2019; Meewan et al., 2022). A recent investigation has proposed the preparation of zein NPs by mixing PEG-modified zein (at least 20%) with unmodified zein through a microfluidics device, thus obtaining stable NPs in physiological medium (van Ballegoie et al., 2022).

According to previous reports, a definite conclusion on zein *in vivo* immunogenicity cannot be driven. More investigations are needed to understand and classify the influence of various factors, such as NP shape, size, surface properties and concentration, drug type and concentration, and administration routes on zein antigenicity.

4 Critical discussion

This review work was focused on the analysis of the physicochemical properties of zein, as a green material for use in nanomedicine. Due to the continuous growth of the world population, the demand for natural resources and the amount of

produced waste materials will increase (Tamasiga et al., 2022). Hence, the implementation of circular economy strategies has become an urgent need. According to the circular economy model, corn waste may be reused as a low-cost raw material for zein extraction, a plant-based protein with a wide range of potential applications in packaging, botany, pharmaceutical and biomedical fields. Zein is low-cost, biocompatible, biodegradable and its amphiphilicity ensures interactions with materials or drugs with different wettability behavior. One interesting zein application explored in this review is its use for the formulation of efficient drug-loaded NPs able to protect therapeutics from early degradation and to convey them to target cells.

To ensure full sustainability, methods for zein NP preparation should also be environmentally friendly. Zein NPs have been mainly produced through phase separation methods, especially using ethanol-water solutions, showing different sizes and drug encapsulation efficiencies depending on solvent types and ratio, zein concentration, zein to drug ratio, mixing type/speed and pH conditions (Figure 3B; Table 3). Typically, the average hydrodynamic sizes of zein NPs were in the 100–400 nm range and drug encapsulation efficiency ranged from about 10% to 85% for both hydrophobic and hydrophilic compounds (Table 3). Despite its acceptable greenness level (Table 2), ethanol replacement with acetone, one of the safest industrial solvents (Joshi and Adhikari, 2019) would enhance sustainability of NP preparation methods. However, to date zein NPs prepared by phase separation using zein solutions in acetone-water have been used for botany applications only (Table 3). The adoption of such methods to the preparation of zein NPs for other applications is thus encouraged.

Sustainable preparation methods for zein NPs would also minimize energy consumption and waste generation (in terms of residual solvents and material). Phase separation methods are advantageous compared to other approaches with high cost and energy consumption, such as electrospraying and supercritical antisolvent precipitation. NP preparation by phase separation, avoiding a subsequent centrifugation step, also minimizes the production of waste material. Furthermore, antisolvent-dialysis combines in one process zein purification from hydrophobic xanthophylls and NP production, decreasing the amount of waste solvent needed for the two consecutive steps.

Although different methods and process conditions have been explored for zein NP preparation (Table 3), one key issue is represented by scale-up production. To this aim, the best zein NP preparation methods are electrospraying, supercritical antisolvent precipitation and microfluidic-based approaches (van Ballegoie et al., 2022). However, among them, only microfluidic methods are environmentally sustainable and low-cost.

Overall, zein NPs are advantageous because they may derive from agricultural waste, as added-value products with efficient drug loading, reproducibility and stability under optimized preparation conditions. Most zein NPs have been applied for the release of lipophilic compounds, such as curcumin, 5-fluorouracil, essential oils, with only a few examples for the release of hydrophilic drugs, such as doxorubicin and chloroquine (Table 3; Table 4). Methods for efficient encapsulation of both hydrophobic and hydrophilic drugs might increase the interest for zein use in nanomedicine. Particularly, gene therapies have recently attracted interest by the scientific community and could represent a promising application for zein NPs. Indeed, to

date only a few studies have reported on zein NPs loaded with plasmid DNAs for gene therapy (Table 3), prepared by pH-triggered complexation. Additionally, such NPs have shown a moderate encapsulation efficiency of 12%–65% (Table 3), deserving improvement. Furthermore, by optimizing preparation methods, zein NPs should be proposed for the encapsulation and release of other oligonucleotide molecules, such as messenger RNAs (mRNAs), microRNAs (miRNAs) and small interference RNAs (siRNAs).

One main application of zein NPs is in oral drug delivery thanks to their resistance to acidic pH, protecting drugs from early degradation. Interestingly, degradation products of zein NPs may also exert beneficial anti-inflammatory effects for their high content of xanthophyll pigments (8%–9%) in yellow zein, such as lutein, zeaxanthin and β -cryptoxanthin. Synergy between zein intrinsic anti-inflammatory effect together with the therapeutic action of encapsulated drugs represents an added-value property.

One of the main reported drawbacks of zein NPs is their poor colloidal stability at physiological conditions. Hybrid core-shell zein NPs have been proposed to address the issue, by the employment polysaccharides, proteins or surfactants as coatings for zein NPs (Table 4). Among them, polysaccharides have been the mostly studied polymers as zein NP coatings, due to their biocompatibility, wide availability and superior stability. Further advantages of core-shell zein-based NPs include higher encapsulation ability and more controlled release kinetics respect to zein NPs. However, in an attempt to propose green zein NP preparation, the addition of other components should not compromise product sustainability. In this regard, zein NPs coated with pectin derived from agricultural waste represent a sustainable solution.

Another key issue of zein NPs is represented by their potential *in vivo* immunogenicity which still remains controversial and would need additional *in vivo* safety studies.

Currently, zein is used as an excipient in food, pharmaceutical industries and vaccine delivery (Tortorella et al., 2021), while drug-loaded zein NPs are still under development and have not reached the approval for clinical use, yet. However, an injectable zein solution, called Ethibloc[®], has been clinically approved in Canada. Ethibloc[®] is percutaneously injected for the treatment of epistaxis, aneurysmal bone cysts, and lymphangiomas (van Ballegoie et al., 2022). Only minor short-term complications related to fever symptoms have been observed, with no long-term side effects. Although Ethibloc[®] is not a nanomedicine product, its approval has demonstrated zein safety for use in humans. Furthermore, one clinical trial is in progress (GLUCOCAPS, Clinical Trial NCT05560412) on the use of zein NPs for glycemic control (Clinica Universidad de Navarra, 2022). According to this trial, patients with prediabetes will consume daily three capsules with nano-encapsulated zein (1 g of zein nano-encapsulated per day), compared to non-encapsulated zein. This study is expected to end in September 2023. Hence, in addition to its higher biocompatibility than synthetic polymers and lipids, and higher safety than animal-derived proteins, zein has an intrinsic therapeutic effect, which might be exploited in synergy with encapsulated drugs. In this regard, more studies are encouraged, particularly on zein use as carrier for oligonucleotide release for the development of new advanced therapies.

Zein NPs have also been investigated for uses in botany for the delivery of pesticides to plants, as an environmental friendly and safer alternative to NPs based on inorganic materials, such as commonly used gold, silver, copper, and iron oxide, which

are affected by overproduction of reactive oxygen species and the release of potentially toxic ions. Finally, based on its biocompatibility and low-cost, the use of zein NPs in veterinary medicine might represent an interesting and not yet explored application field.

5 Conclusion

Zein has been proposed as a biocompatible and biodegradable plant-based protein for drug delivery applications. Extensive literature on the topic showed that zein NPs are advantageous because of the availability and low cost of the raw material. Physicochemical properties of zein such as aminoacidic composition and solvent miscibility have been demonstrated to be critical parameters for the preparation of green zein NPs. Among the available production methods, we underlined superiority of phase separation methods using acetone-water solutions for a sustainable preparation of drug-loaded zein NPs. Antisolvent-dialysis method was also found to be an interesting option for zein NP preparation, as able to combine zein purification and NP production in one step, thus decreasing the volumes of used solvent. On the other hand, zein NP production scalability is an important issue for lab to market translation. In this regards, novel microfluidic based approaches, based on phase separation methods, represent an optimal and sustainable choice.

Zein NPs have been mainly employed to deliver lipophilic drugs, such as curcumin and essential oils: such drugs are efficiently encapsulated by zein NPs because they interact with the hydrophobic aminoacidic residues of zein. However, a few recent investigations have also highlighted the possibility to release hydrophilic compounds from zein NPs, among which oligonucleotides. In this regard, examples reported so far have only focused on the encapsulation of plasmid DNAs into zein NPs, while the loading of other oligonucleotides such as mRNAs, siRNAs and miRNAs has yet to be investigated and could pave the way to the application of zein NPs for advanced RNA-based therapies.

To conclude, this review provides an innovative understanding of zein NP state of the art and provides useful information for the development of a new generation of sustainable, biodegradable, safe and innovative drug delivery vectors based on zein.

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Author contributions

FT: conceptualization, formal analysis, writing—original draft; VC: supervision (project supervision); conceptualization, writing—review and editing, project administration, resources, funding acquisition. All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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