Oregano Essential Oil Micro- and Nanoencapsulation With Bioactive Properties for Biotechnological and Biomedical Applications

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Due to the preservative, antioxidant, antimicrobial, and therapeutic properties of oregano essential oil (OEO), it has received an emerging interest for biotechnological and biomedical applications. However, stability and bioactivity can be compromised by its natural volatile and hydrophobic nature, and by external factors including light, heat, or oxygen. Therefore, micro- and nanoencapsulation are being employed to guarantee oregano oil protection from outside aggressions and to maximize its potential. Oregano oil encapsulation is an interesting strategy used to increase its stability, enhance its bioactivity, and decrease its volatility. At the same time, the versatility that micro- and nanocarriers offer, allows to prepare tailored systems that can provide a controlled and targeted release of the encapsulated principle, influence its bioactive activities, or even provide additional properties. Most common materials used to prepare these carriers are based on lipids and cyclodextrins, due to their hydrophobic nature, polymers due to their versatility in composition, and hybrid lipid-polymer systems. In this context, recently developed micro- and nanocarriers encapsulating oregano oil with applications in the biotechnological and biomedical fields will be discussed.

Keywords: microcarriers, nanocarriers, antimicrobial, antibacterial, antifungal, emulsion, stability

INTRODUCTION

Essential oils (EOs), derived from aromatic plants, are volatile oily liquids mainly composed of terpenoids and phenolic acids (da Silva et al., 2021). They have been used since ancient times in different cultures due to their bioactive properties. Some of the most reported properties of EOs are their antibacterial (Nazzaro et al., 2013), antifungal (D’agostino et al., 2019), antiviral (Ma and Yao, 2020), and antioxidant (Leyva-López et al., 2017) activities, mainly due to the disruption of bacterial and fungal membranes and viral envelopes (Böhme et al., 2014). Nevertheless, some characteristics like immunomodulatory and anticancer activities are recently being reported, highlighting the potential use of EOs in the biomedical field (Bhalá et al., 2013; Böhme et al., 2014). For these reasons, in the last decades, it has emerged a great interest in their use in biotechnology, for example, in foods and cosmetics, and in the biomedical field, in which their excellent properties provide a great therapeutic potential (Böhme et al., 2014; Aljaafari et al., 2021). However, stability and bioactivity can be compromised by their natural volatility, low water solubility, and external...
factors, which have detrimental effects on the overall acceptability of the developed product (Turek and Stintzing, 2013).

In this context, drug delivery systems such as micro- and nanoparticles (MPs and NPs), micro- and nanocapsules (MCs and NCs), films, or nanocomposite materials have been proposed to encapsulate EOs (Zhu et al., 2021). These systems enhance EO stability in aqueous media and, as a consequence, improve their bioavailability, reduce their toxic effects, provide a controlled release of the encapsulated agent, protect them against the environment or mask their intense aroma (Cimino et al., 2021). Micro- and nanocarriers with tailored properties are of special interest due to the increased surface-to-volume ratio that their sizes offer and, consequently, an increase in their reactivity (Franklyne et al., 2016). These systems are typically based on polymers, lipid materials, or a combination of both (Kaliamurthi et al., 2019). Moreover, micro- and nanocarriers present some differences regarding their fate after its application, the ability to cross some biological barriers, entering cells, and possible tissue reactions that, depending on the application, will determine the choice of one over the other (Kohane, 2007).

Oregano essential oil (OEO) is one of the most widely used EOs worldwide. It is extracted from *Origanum vulgare* L. and formed basically by carvacrol and thymol (Teixeira et al., 2013). Both carvacrol and thymol are monoterpenes with a single phenolic ring formed from the bonding of two isoprene molecules with three functional group substituents (Memar et al., 2017). Due to this chemical structure, they provide OEO with its antibacterial and antioxidant properties, in addition to its anticancer and anti-inflammatory activities (Sakkas and Papadopoulou, 2017; Sharifi-Rad et al., 2021). Due to these activities, OEO and its components have come to the forefront and are being widely investigated to be used as a food preservative, for active packaging, and the treatment of different diseases, such as infections (Bhalla et al., 2013).

This review aims to discuss the state-of-the-art micro- and nanoencapsulation of OEO in biotechnology and biomedical applications, making emphasis on the materials used, the fabrication process, and their final bioactive properties. It is expected that the information provided here will provide the reader with a general view of the possibilities that OEO encapsulation may offer in these specific fields. In this sense, Figure 1 collects the main types of OEO delivery systems reported, their fabrication methods, most common materials employed, and biomedical and biotechnological applications.

**REVIEW METHOD**

For this mini-review article, an extensive search was conducted in different web search engines using keywords such as OEO, carvacrol, thymol, microencapsulation, nanoencapsulation, microcarriers, nanocarriers, MPs, NPs, MCs, NCs, biomedical and biotechnological. The search strategy was limited to publications in English and published from 2017 to 2021. Articles were classified according to the size of the OEO delivery system and listed according to their application.

**OEO MICROENCAPSULATION**

Microencapsulation is a technique in which a material of interest is surrounded by a coating to form capsules or
particles with sizes between 1 and 1,000 µm (Ju and Chu, 2019). Compared to macroscale particles, MPs and MCs have the advantage of having a larger surface-to-volume ratio, which is even larger for NPs, increasing the reactivity of the delivery system. Microencapsulation of OEO has been conducted through different methods such as emulsification, spray-drying, coaxial electrospray, freeze-drying, coacervation, \textit{in situ} polymerization, or ionic gelation using mainly lipids, cyclodextrins, and polymers (Bakry et al., 2016). One of the main applications of these OEO-loaded systems is in the field of biotechnology as food preservatives, as components of active packages, and in the pharmaceutical industry (Bakry et al., 2016). Research articles published on this topic in the latest years and the main results obtained are listed in Table 1 according to the system function or application.

The stability of microemulsions, MPs, and MCs encapsulating OEO has been widely investigated. In this sense, Cardoso-Ugarte et al. (2021) optimized OEO loaded water-in-oil-in-water (W/O/W) double emulsions in terms of primary emulsion concentrations and homogenization parameters to finally assess the antifungal activity of the best system (Cardoso-Ugarte et al., 2021). They assessed the stability of primary emulsions resulting from high-pressure and mechanical homogenization evidencing that stability, in terms of droplet size, was affected by the homogenization method of the primary emulsion. Asensio et al. (2017), for their part, studied how wall materials (hydroxypropyl methylcellulose, maltodextrin, and colloidal silicon dioxide) and storage temperatures influenced the antioxidant activity, total phenolic content, and release kinetics of OEO compounds of spray-dried microcapsules (MCs) (Asensio et al., 2017). The authors observed that the presence...
of colloidal silicon dioxide in some formulations increased the release of volatile compounds, to detriment of the antioxidant activity, in terms of radical scavenging activity and Trolox equivalent antioxidant capacity, which was lost to a greater extent during storage.

Highly stable Pickering emulsions (emulsion stabilized by solid particles) loading OEO were prepared by Zhou et al. (2018) using cellulose nanocrystals as stabilizers (Zhou et al., 2018). The authors demonstrated that emulsions exhibited higher stability with increasing concentration of cellulose nanocrystals or at lower oil-water ratio and salt concentration. Furthermore, the antimicrobial efficacy of the emulsions was confirmed by efficiently inhibiting the growth of different bacteria by destroying the integrity of their cell membranes.

### TABLE 1 | Latest research reports (2017–2021) on oregano essential oil (OEO) microencapsulation with biotechnological and biomedical applications.

| Function/application | Fabrication process | Composition | Results | References |
|----------------------|---------------------|-------------|---------|------------|
| Delivery system      | W/O emulsion        | OEO, corn oil | Emulsifier concentration affected emulsion separation, viscosity, and surface electric charge. | Cardoso-Ugarte et al., 2018 |
| Delivery system      | Emulsion            | Thymol, alginate | No chemical interactions between thymol and sodium alginate. Sustained in vitro release. | Bhalerao and Wagh, 2019 |
| Antifungal           | W/O/W double emulsion | OEO, corn oil | Stability and droplet sizes are affected by the homogenization method of the primary emulsion. Antifungal activity against A. niger. | Cardoso-Ugarte et al., 2021 |
| Antimicrobial agent   | Pickering emulsion  | OEO, cellulose nanocrystals | Average droplet sizes: 1.2–2.9 µm. Variations of zeta potential with pH. Antimicrobial activity by cell membrane disruption. | Zhou et al., 2018 |
| Antimicrobial agent   | O/W emulsion        | OEO, high oleic sunflower oil | Nanemulsions were not stable under acid and high salt concentration conditions. Inutec SP1 emulsions remained stable for several days. | Sedaghat Doost et al., 2017 |
| Antimicrobial agent   | Solvent evaporation | Thymol, PLGA | Improvement of thermal and storage stability sustained release of thymol and antibacterial properties against E. coli and S. aureus. | Zhu et al., 2019 |
| Antimicrobial agent   | Spray drying        | Carvacrol, pectin-alginate | Encapsulation efficiency: 77%. Antimicrobial activity against E. coli. | Sun et al., 2019 |
| Antioxidant additive | Spray drying        | OEO, hydroxypropyl methyl cellulose, maltodextrin, colloidal silicon dioxide | Release kinetics of OEO volatile compounds and the antioxidant activity of the system was controlled by wall material-lipid core ratios and temperature storage conditions. | Asensio et al., 2017 |
| Food preservation    | Co-precipitation inclusion method | OEO, β-cyclodextrin | Inclusion efficiency: 55%. Enhancement of yam shelf-life. | Huang et al., 2020 |
| Food preservation    | Ionic gelation and electrostatic interactions (coating) | OEO, alginate, whey protein concentrate | Loading and retention capacity: 64 and 57%. Freeze-drying caused an increase in size polydispersity and porosity of bead surface. Whey protein coating caused a slowdown effect on OEO release. | Gallo et al., 2020 |
| Active packaging     | Co-precipitation inclusion method | OEO, β-cyclodextrin | Particle size range: 450–530 nm. Polydispersity: 0.31–0.48. Good stability in suspension. Inclusion efficiencies: 26%. In vitro OEO release for up to 11 days. | Kotronia et al., 2017 |
| Topical treatment of cutaneous diseases | Double emulsion | OEO | Average particle size: 1.76 µm. Surface charge: −15 mV. Encapsulation efficiencies: 47%. Unstable with temperature. | Fraj et al., 2019 |
| Topical cosmetics    | Ultrasonication     | Thymol, lignosulfonate | Average particle size range: 3.2–3.4 µm. Encapsulation efficiencies >40%. In vitro slow release of thymol using simulated fluids. | Piombino et al., 2020 |
| Antibiotic substitute for intestinal delivery | Melt-granulation process | Thymol, lauric acid, starch, alginate | Stability during feed pelleting and storing processes. In vitro and in vivo prolonged release of thymol in simulated gastric and intestinal fluids and pig guts, respectively. | Omonjio et al., 2018 |
| Intestinal delivery  | Blending of thymol with a lipid matrix | Thymol, commercial lipid matrix containing other organic acids | Stability during feed pelleting and storing processes. In vitro and in vivo prolonged release of thymol in simulated gastric and intestinal fluids and pig guts, respectively. | Choi et al., 2020 |

W/O, water-in-oil; W/O/W, water-in-oil-in-water; O/W, oil-in-water; PCL, polycaprolactone; PLGA, poly(lactic-co-glycolide).
The influence of environmental stress conditions, such as acidification and salt addition, on the stability of OEO oil-in-water (O/W) emulsions prepared by a high energy method, has also been assessed by Sedaghat Doost et al. (2017), using two non-ionic surfactants, Tween 80 and Inutec SPI, as stabilizers (Sedaghat Doost et al., 2017). Different OEO: high oleic sunflower oil ratios were used to prepare the emulsions. Despite nanoemulsions could be formed, they were not stable under acid and high salt concentration conditions. Moreover, Tween 80 containing emulsions exhibited phase separation at all salt concentrations, while Inutec SPI emulsions remained stable for several days. In the end, colloidal dispersions with a 50:50 OEO: high oleic sunflower oil ratio in the lipid phase stabilized by Inutec SPI kept at 4°C showed the longest stability with no droplet size variation during 2 weeks. The effect of Tween 80 in addition to Span 20 as emulsifiers of OEO water-in-oil (W/O) emulsions was also investigated (Cardoso-Ugarte et al., 2018). In this case, the concentration of the emulsifier affected emulsion separation, led by Ostwald ripening, viscosity, and surface electric charge, showing slower separation rates in the tested emulsions with higher concentrations of Tween 80 and Span 20.

Cyclodextrins, cyclic hydrophillic oligosaccharides obtained from starch enzymatic conversion, have been employed in OEO encapsulation. For instance, Kotronia et al. (2017) encapsulated OEO into β-cyclodextrin inclusion complexes by co-precipitation methods (Kotronia et al., 2017). The systems showed suitable characteristics in terms of size, surface charge, and morphology, a controlled in vitro OEO release for up to 11 days, and inclusion efficiencies up to 26%. Similarly, Huang et al. (2020) prepared β-cyclodextrin systems loaded with OEO, obtaining MPs with strong interactions between β-cyclodextrin and OEO. A reduction of both gram-positive and gram-negative bacteria occurred when treated with OEO-loaded systems, especially in gram-positive bacteria. Finally, the food preservative performance of the system due to its antibacterial activity was demonstrated by reducing the browning and enhancing the shelf-life of a type of yam (Huang et al., 2020).

Alginate beads have also been used to encapsulate OEO by extrusion dripping to study the influence of its encapsulation in oil release kinetics in liquid simulating meat marinating solution (Gallo et al., 2020). Spherical particles with good encapsulation performance were obtained and subjected to electrostatic interactions with whey proteins and freeze-drying. Freeze-drying of the beads increased the particle size polydispersity and the porosity of the bead surface. Besides, beads were successfully coated with whey proteins by electrostatic interactions causing a slowdown effect on OEO in vitro release rates. Alginate has also been used to encapsulate thymol, one of the main components of OEO, demonstrating that no chemical interactions between thymol and sodium alginate occurred and showing sustained thymol in vitro release (Bhalerao and Wagh, 2019). Carvacrol, the other main component of OEO has also been microencapsulated, in a pectin-alginate matrix by Sun et al. (2019), demonstrating that the microencapsulation did not affect the radical scavenging properties of carvacrol and the antibacterial activity against Escherichia coli (Sun et al., 2019).

Thymol has been entrapped in other lipid and polymeric vehicles. For instance, Omonjio et al. (2018) microencapsulated thymol along with lauric acid into starch MPs with the presence or not of alginate to deliver them to pig intestinal tracts as an antibiotic substitute (Omonjio et al., 2018). Highly stable systems were obtained with an in vitro prolonged release of loaded compounds in the case of MPs presenting alginate, using simulated salivary, gastric, and intestinal fluids. However, the efficacy of the MPs was not demonstrated. The intestinal delivery of thymol has also been assessed when microencapsulated into commercial lipid matrices containing organic acids by Choi et al. (2020), demonstrating the stability of the systems during feed pelleting and storing processes and an in vitro and in vivo sustained release of thymol in simulated gastric and intestinal fluids and pig guts, respectively (Choi et al., 2020).

Due to the antioxidant and biocompatibility properties of lignin, lignosulfonate MPs were successfully developed by Piombino et al. (2020) to encapsulate thymol and its derivatives as topical systems with antimicrobial properties for cosmetics through environmental friendly sonication procedures. Results showed that more than 40% of each substrate was properly encapsulated, except for 2,4-dibromothymol, showing the best encapsulation efficiencies for the mono-brominated thymol derivative (76%). To test the suitability of these systems as dermal agents, the in vitro release of the derivatives from the MPs in solutions simulating skin pH (acetate buffer at pH 5.4) were performed, showing a slow-release, especially for O-methylated compounds, that was dependent on the inherent lipophilicity of each compound. Thymol-loaded poly(lactic-co-glycolide) (PLGA) MPs have also been demonstrated to be suitable microcarriers of thymol improving thermal and storage stability and controlling thymol release (Zhu et al., 2019). In addition, the antibacterial properties of the MPs against E. coli and Staphylococcus aureus were demonstrated by the disruption of their cytoplasmic membrane, since the porous structure of the MPs enhances the permeation of thymol into bacteria. The antibacterial effect was confirmed by adding the loaded MPs into naturally contaminated milk and observing that the growth of bacteria was suppressed by thymol-loaded MPs.

**OEO NANOENCAPSULATION**

Nanoencapsulation is another common strategy addressed to protect OEO from the environment and improve its performance (Bilia et al., 2014). In addition, the smaller size of NPs, between 1 and 1000 nm, makes them very suitable for biomedical applications, as they can be intravenously injected, can be efficiently uptaken by a variety of cell types, and can extravasate through endothelium to reach, for instance, inflammatory sites or tumors (Gelperina et al., 2005; Singh and Lillard, 2009). The latest research reports on OEO nanoencapsulation that are mainly focused on food biotechnology and biomedical applications are listed in Table 2 according to the system function or application.

Polymeric nanocarriers have received great interest due to their versatility in composition, structure, and properties. In the study of Fraj et al. (2019), the authors compared
TABLE 2 | Latest research reports (2017–2021) on OEO nanoencapsulation with biotechnological and biomedical applications.

| Function/application | Fabrication process | Composition | Results | References |
|----------------------|---------------------|-------------|---------|------------|
| Delivery system      | High energy emulsion method | OEO, sunflower oil, succinic anhydride-modified starch, chitosan, sodium carboxymethylcellulose | Multilayer NPs: One layer NPs: 180 nm, −42 mV. Two layers NPs: 226 nm, 35 mV. Three layers NPs: 265 nm, −1 mV. Encapsulation efficiency: 97%. | Espinosa-Sandoval et al., 2021 |
| Delivery system      | Complex coacervation | OEO, gelatin, chia mucilage/arabic gum | Particle size range: 17–120 nm. Encapsulation efficiency: >96%. | Hernández-Nava et al., 2020 |
| Delivery system      | Ultrasonication | Neobee® 1053 medium-chain triglyceride oil | Lecithin nanoemulsions were more stable and viscoelastic than Tween 20 nanoemulsions. | Nash and Erk, 2017 |
| Antifungal agent for food preservation | Electrospraying | OEO, PVA, chitosan | Particle size range: 337–818 nm. Encapsulation efficiency: 90%. | Vehipi et al., 2020 |
| Antimicrobial agent for food preservation | Emulsion | OEO, medium-chain triacylglyceride | Average droplet size range: 74–150 nm. Viscoelastic behavior of nanoemulsions. Antimicrobial activity by quorum-sensing inhibition. | Asensio et al., 2020 |
| Antimicrobial agent for food preservation | High-frequency ultrasonication | OEOs (cinnamon, rosemary, OEO) | Particle size range: 226–546 nm. Encapsulation efficiency >80%. | Dávila-Rodríguez et al., 2019 |
| Antibacterial agent for food preservation | Nanoparticles: lipid film hydration technique | Carvacrol, soy phosphatidylcholine | Nanoparticles: Particle size: 271 nm. Zeta potential: 8.6 mV. Encapsulation efficiency: 98%. | Ayres Cacciatore et al., 2020 |
| Antibacterial agent for food preservation | Nanocapsules: interfacial deposition technique | Carvacrol, Eudragit® | Particle size: 159 nm. Zeta potential: 44.8 mV. Encapsulation efficiency: 97%. | Nash and Erk, 2017 |
| Antibacterial agent for food preservation | Standard Schlenk techniques by ring-opening polymerization | PEI, PLA | Average particle size: 115 nm. Zeta potential: 55 mV. Encapsulation efficiency: 54%. Enhanced antibacterial effect. | Niza et al., 2020 |
| Antimicrobial agent | Single emulsion | Thymol, PLA | Average particle size range: 220–260 nm. Encapsulation efficiency: 60%. Improved antibacterial effect compared to non-encapsulated thymol. | Marcet et al., 2018 |
| Antimicrobial agent | Novel, simple chemical synthesis | Thymol, chitosan, silver | Average particle size: 29 nm. Spherical shape, monodisperse in water, excellent blood biocompatibility. | Manukumar et al., 2017 |
| Topical treatment of cutaneous diseases | Nanoprecipitation | OEO, PCL | Average particle size: 181 nm Polydispersity: 0.133. Surface charge: −41 mV. Encapsulation efficiencies: 85%. | Fraj et al., 2019 |
| Anti-angiogenic system | Ultrasonication | Carvacrol, medium chain triglyceride | Hydrodynamic droplet size: 101 nm. Zeta potential: −39 mV. Decrease in the expression of several angiogenic markers in a lung adenocarcinoma model. | Khan et al., 2019 |
| Pharmaceutical product for airway lung disease | Fusion-emulsification | Carvacrol, cocoa butter, 3.5-dii-tet 4-butylhydroxytoluene, imidazolidinyl urea | Minimization of oxidative stress and histological damage generated from smoke inhalation in rodents. | Carvalho et al., 2020 |

OEO, oregano essential oil; PVA, polyvinyl alcohol; EO, essential oil; NP, nanoparticle; PEI, polyethylenimine; PLA, polylactic acid; PCL, polycaprolactone.

the properties of OEO-loaded NPs and MPs prepared by nanoprecipitation and double emulsion, respectively (Fraj et al., 2019). Results demonstrated that, while NPs were stable at different temperatures, MPs suffered an increase in particle size and a decrease in carvacrol component retention. In other studies, gelatin combined with chia mucilage was used as an alternative OEO delivery system to other most commonly used in complex coacervation (gelatin combined with Arabic gum).
NPs of both gelatin and chia mucilage and gelatin and Arabic gum were synthesized obtaining high encapsulation efficiencies in both cases. Moreover, the amount of Tween 80 and OEO concentration influenced the NP size, obtaining smaller particles with increasing Tween 80 due to the enhancement of the interfacial tension reduction and droplet breaking. Finally, complex coacervates with the highest encapsulation efficiencies were spray-dried obtaining the best flow properties for the gelatin-chia mucilage NPs.

Chitosan was used by Espinosa-Sandoval et al. (2021) to prepare OEO-loaded multilayer nano-emulsions by high energy methods (Espinosa-Sandoval et al., 2021). In this study, octenyl succinic anhydride-modified starch combined with partially deacetylated chitosan of medium and low molecular weight and carboxymethylcellulose was used to protect OEO in a multilayer system. Interestingly, using an in vitro gastric condition simulating test, the authors showed that each polymeric layer influenced OEO bioaccessibility, obtaining the highest value for the three-layer system. Spherical polymeric NPs based on polyvinyl alcohol (PVA) and chitosan loaded with OEO have also been prepared by electrospaying (Vehapi et al., 2020). Results demonstrated that the encapsulation of OEO and the presence of chitosan led to a superior antifungal effect of the nanoeencapsulated system compared to free OEO.

Oregano essential oil nanoemulsions have also been reported to obtain nanocarriers with antibacterial properties. For instance, Asensio et al. (2020) prepared OEO nanoemulsions, showing that the incorporation of the oil could increase nanoemulsion stability, lower droplet size, and increase emulsion viscosity (Asensio et al., 2020). Furthermore, nanoemulsions exhibited good antimicrobial activity by the inhibition of cell-to-cell communication of gram-negative bacteria, also known as quorum-sensing (ability to detect and respond to cell population density by gene regulation). Dávila-Rodríguez et al. (2019) prepared O/W nanoemulsions encapsulating three different EOs, cinnamon, rosemary, and OEO, using a high-frequency ultrasound technique. Results proved that nanoencapsulated EOs were more effective than free EOs since a lower amount of EO was required to provide the antibacterial effect. Moreover, OEO nanoemulsions proved to be the most effective antimicrobial systems against E. coli and Listeria monocytogenes (Dávila-Rodríguez et al., 2019).

Another advantage that encapsulation can offer is masking the intense aroma of OEO components. Carvacrol encapsulation into nanostructures, nanoliposomes, and polymeric Eudragit® NCs were developed by Ayres Cacciatore et al. (2020), establishing that its encapsulation could be interesting to reduce its aroma due to its controlled release (Ayres Cacciatore et al., 2020). The effect of lecithin or Tween 20 on O/W nanoemulsions encapsulating carvacrol was studied by Nash and Erk (2017), concluding that, while lecithin nanoemulsions were highly viscoelastic and gave stability to the nanoemulsion, Tween 20 did not (Nash and Erk, 2017).

Oregano essential oil encapsulation using polycationic polymers can improve its antibacterial effect due to the enhanced bacterial uptake that positive charges cause. Polyethylenimine (PEI)-coated polylactic acid (PLA) NPs encapsulating carvacrol have also been developed as antimicrobial agents (Niza et al., 2020). In this study, NPs coated with PEI possessed a positively charged surface that facilitated their uptake by bacteria compared to negatively charged ones, enhancing the antibacterial effect of the encapsulated carvacrol. Moreover, NPs displayed higher antibacterial activity than free carvacrol, a sustained release, and stability during storage.

Other biomedical applications of carvacrol-loaded nanocarriers have been reported. Khan et al. (2019) demonstrated the potent anti-angiogenic effect both, in vitro and in vivo, of carvacrol-loaded O/W nanoemulsions, by reducing the expression of several angiogenic markers such as COX-2, VEGF, and CD31 in a lung adenocarcinoma model (Khan et al., 2019). Furthermore, solid lipid NPs incorporating carvacrol were prepared to treat lung damage of airway smoke inhalation and tested in an in vivo rat model by Carvalho et al. (2020). Results showed that carvacrol-loaded solid lipid NPs could minimize oxidative stress and inhalation injury and histological damage generated from smoke inhalation in rodents compared to the negative control.

To improve the antibacterial effect of thymol, different nanocarriers have been developed. An example is that of biodegradable PLA NPs developed by Marcet et al. (2018). PLA was found to be the key variable in optimizing NP preparation in terms of size and encapsulation efficiency, producing NPs with high storage stability at several pHs and improved antimicrobial properties compared to non-encapsulated thymol. In other studies, thymol was loaded into intrinsic antibacterial chitosan silver NPs showing interesting antioxidant properties through radical scavenging and antibacterial activity due to the three components against different gram-positive bacterial strains (Manukumar et al., 2017).

**CONCLUSIONS**

Although oregano oil has been used in different cultures since ancient times, it has received special attention in the last decades due to its preservative, antimicrobial, and therapeutic characteristics. However, its bioactivity is compromised by its highly volatile and hydrophobic nature and by external environmental factors. Hence, different drug delivery systems are being explored as a strategy to increase its stability and bioavailability, protect it from the environment, control its release, and even enhance its properties. In this review, the latest research on micro- and nanocarriers encapsulating OEO focusing on biomedical and biotechnological applications revealed that carriers such as emulsions and polymeric-based systems seem to be the most appropriate ones for encapsulation of this compound. Stability has been demonstrated to be particularly important in this type of delivery system since it determines the final performance of the loaded system, and it is dependent on multiple parameters such as the composition, the fabrication method, and the storing conditions. On the contrary, the mechanism of action by which OEO exerts its activities is not deeply investigated in most of the reviewed articles, but instead, they focused on the properties and final performance of the loaded delivery system. Overall, the process of OEO encapsulation stands out as a possible alternative for the preservation of this oil against environmental conditions,
increasing its stability and maintaining its bioactive properties, mainly its antioxidant and antimicrobial ones. Furthermore, design and formulation on the carrier employed to encapsulate this oil, can influence and enhance its bioactive properties and even can provide the final delivery system with additional and beneficial properties.

AUTHOR CONTRIBUTIONS

GP-Q: conceptualization and writing—original draft. SE-R: conceptualization, project administration, and funding acquisition. JP: conceptualization, resources, and funding acquisition. MA and BV-L: conceptualization, resources, writing—review and editing, supervision, project administration, and funding acquisition. All authors contributed to the article and approved the submitted version.

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Conflict of Interest: JP is the CEO and founder of Alodia Farmacéutica SL and SE-R is an employee of the same company.

The remaining 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.