Strategic developments in the drug delivery of natural product dihydromyricetin: applications, prospects, and challenges

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ABSTRACT
Dihydromyricetin (DHM) is an important natural flavonoid that has attracted much attention because of its various functions such as protecting the cardiovascular system and liver, treating cancer and neurodegenerative diseases, and anti-inflammation effect, etc. Despite its great development potential in pharmacy, DHM has some problems in pharmaceutical applications such as low solubility, permeability, and stability. To settle these issues, extensive research has been carried out on its physicochemical properties and dosage forms to produce all kinds of DHM preparations in the past ten years. In addition, the combined use of DHM with other drugs is a promising strategy to expand the application of DHM. However, although invention patents for DHM preparations have been issued in several countries, the current transformation of DHM research results into market products is insufficient. To date, there is still a lack of deep research into the pharmacokinetics, pharmacodynamics, toxicology, and action mechanism of DHM preparations. Besides, preparations for combined therapy of DHM with other drugs are scarcely reported, which necessitates the development of dosage forms for this application. Apart from medicine, the development of DHM in the food industry is also of great potential. Due to its multiple effects and excellent safety, DHM preparations can be developed for functional drinks and foods. Through this review, we hope to draw more attention to the development potential of DHM and the above challenges and provide valuable references for the research and development of other natural products with a similar structure-activity relationship to this drug.

GRAPHICAL ABSTRACT

1. Introduction
Dihydromyricetin (2R,3R-3,5,7,3′,4′,5′-hexahydroxy-2,3-dihydroflavonol, DHM) is a polyphenol hydroxyl dihydroflavonol compound (Figure 1A and 1B) with high content in ampelopsis grossedentata which has long been used for herbal tea and traditional Chinese medicine in China (CC Sun et al., 2021). Many studies have shown that DHM has many pharmacological activities. For example, DHM can protect the cardiovascular system through PI3K/Akt, Nrf2/ARE, and SIRT3 signal pathways, and can also prevent or inhibit gastric cancer, lung cancer, hepatocellular carcinoma, colorectal cancer, breast cancer, and melanoma through Akt/STAT3, ERK/Akt, Akt/Bad, AMPK/AMPK/XAF1, Akt-mTOR, ROS/NF-κB, and mitochondrial apoptosis signal pathways (Liu et al., 2016; KJ Fan et al., 2017; Park et al., 2017; Zhang et al., 2017; Zhou et al., 2017; Chen et al., 2018; Li et al., 2021; Wang et al., 2021; Zhang et al., 2021). DHM can treat neurodegenerative diseases by Nrf2, SIRT1, and p53/p21 signal pathways, and can improve alcoholic liver disease, alcoholic fatty liver disease, and acute liver injury through JNK/AP-1/iNOS, p53, TGF-β1/Smad3, and AMPK/PGC-1α/ERRα.

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signal pathways (Xie et al., 2015; Kou et al., 2016; Hu et al., 2018; JQ Ma et al., 2019; Yan et al., 2019; Zeng et al., 2019). DHM also has anti-inflammatory, antibacterial, antiviral, and skin protection effects, so it has great development potential (Figure 1C) (Moon et al., 2018; Liang et al., 2020; Sun et al., 2021; Landis et al., 2022; Shi et al., 2022). DHM has not been reported to be toxic in the conventional dose range, and there is no significant change in physiological indexes such as the body weight and temperature after oral administration of 150–1500 mg/kg DHM in rats (Ma et al., 2014). Moreover, no obvious cytotoxicity of DHM has been found in normal cells, which indicates its high bio-safety (Cai et al., 2016).

DHM has poor water solubility and only exists stably in low temperatures and weak acidic environments (pH 6.0) (CC Sun et al., 2021). If the temperature exceeds 100 °C, an irreversible oxidation reaction could occur. At 25 °C and 37 °C, the solubility of DHM in water is about 0.2 mg/ml and 0.9 mg/ml, respectively (L Fan et al., 2017). To increase its solubility, permeability, and stability, researchers have developed many new dosage forms for DHM, such as nano-preparations, microemulsions, gels, crystals, phospholipid complexes, and cyclodextrin complexes, etc., thereby improving the medicinal properties of DHM (CC Sun et al., 2020; S Geng, Jiang, et al., 2021; L Liu et al., 2022). It has been reported that DHM can distribute in various tissues and can pass through the blood-brain barrier after oral administration in rats, which provides further conditions for curing various diseases (L Fan et al., 2017). In addition, since the effect of this drug is not ideal when used alone because of its low titer, synergistic application of DHM with other anti-tumor drugs, liver protection drugs, anti-neurodegenerative diseases drugs, etc., should be considered.

So far, the physicochemical properties, analytical methods, pharmacological properties, and potential mechanisms of DHM have been reviewed in detail elsewhere (D Liu et al., 2019; Martinez-Coria et al., 2019; CC Sun et al., 2021). Sun et al. fully reviewed the physicochemical properties of DHM and discussed the effect of ascorbic acid on its stability and bioavailability (CC Sun et al., 2021). In addition, Liu et al. detailed the quantification methods and the main biological characteristics of DHM, as well as the methods to enhance its bioavailability (D Liu et al., 2019). Moreover, the pharmacological research progress and potential mechanisms of DHM in anti-tumor, metabolic diseases, liver diseases, cardiovascular diseases, skin diseases, and neurological diseases have also been reviewed in detail (Z Zhang et al., 2018; Martinez-Coria et al., 2019; Tong et al., 2020; J Chen et al., 2021; L Chen et al., 2021; Wu et al., 2022). Therefore, these contents will not be reiterated here. As DHM has poor water solubility, permeability, and stability, over the past decade, many modern techniques have been adopted to design various dosage forms for DHM to settle these problems. However, although previous studies have reported a large number of DHM preparations, there is a lack of attention to the problems in the application of these preparations. Furthermore, despite the potential dosage forms reported in the previous articles and patents, most of these studies only focused on the preparation and the characterization of DHM preparations but lacked further investigations such as toxicology studies and drug efficacy studies that are essential for the transformation of this drug into clinical applications. Moreover, there are many other issues to consider in the future application of DHM. Based on the facts above, this review focused on the pharmaceutical designs of DHM and analyzed the reported dosage forms and some potential dosage forms of DHM. Through this review, we also analyzed the challenges and potential strategies of pharmaceutical design for the future applications of this drug combined with other drugs (e.g. antitumor drugs, antiviral drugs, antibacterial drugs, and hepatoprotective drugs), as well as some other issues to consider in the drug delivery of DHM, in order to provide valuable references for the development and application of DHM in the medicine and food industries and provide ideas.
for the research and development of natural products with a similar structure to DHM.

2. Pharmaceutical designs of DHM

2.1. Nano-preparations

2.1.1. Liposomes

Liposomes are drug carriers with the advantages of low toxicity, high biocompatibility, and low immunogenicity, which can carry and contain various hydrophilic and lipophilic drugs (Yuba, 2020). Common liposomes are easily engulfed by the mononuclear phagocyte system after entering the blood as drug carriers, leading to their rapid clearance (Cheng et al., 2022). To overcome such problems, modification of liposomes (e.g. PE Gylation) is usually adopted to reduce the immunogenicity and improve the stability and the circulation time of liposomes (Ibaraki et al., 2021). Luo et al. used egg yolk phospholipids and cholesterol as carrier materials to prepare tea saponin (TS) coated polyethylene glycol DHM nanoliposomes (TS/CTS@DHM-lips). In these nanoliposomes chitosan (CTS) and TS form a cationic coating, which made the liposome surface positively charged, thus showing deep cell membrane permeability to negatively charged Escherichia coli and Staphylococcus aureus by electrostatic interaction, while DHM exerted increased anti-bacterial effect via sustained release through the coating. TS/CTS@DHM-lips had excellent entrapment efficiency and drug loading, and improved the hydrophilicity and solubility of DHM, making it a potential antibacterial agent (F Luo, Zeng, Chen, et al., 2021). In another study, DHM nanoliposomes prepared by Zhang et al. with soybean phosphatidylcholine and cholesterol as carrier materials and distearoyl phosphatidylethanolamine-polyethylene glycol-2000 as modifier reduced the release rate of DHM in vitro and prolong the residence time of DHM in rats, which improved the oral bioavailability of DHM (W Zhang et al., 2018). However, repeated intravenous injection of PE Gyalted liposomes into the same animal has been reported to result in an accelerated blood clearance phenomenon (Dams et al., 2000; El Sayed et al., 2020). Therefore, it is urgent and necessary to develop a strategy that can reduce or eliminate the immune response induced by PE Gyalted liposomes without affecting the therapeutic effect in vivo.

Multivesicular liposomes (MVL) are aggregates formed by close packing of non-concentric lipid bimolecular vesicles. Compared with ordinary liposomes, MVL can achieve high encapsulation and low leakage of drugs because of its ‘foam’ structural characteristics and ensure the storage and release of drugs in their original forms (Chaurasiya et al., 2022). DHM liposomes (DHM-lips) prepared by Luo et al. with cholesterol and egg yolk lecithin as drug carriers and PEG-4000 as modifier could effectively improve the solubility and membrane permeability of DHM, prolong the release rate and anti-bacterial time of DHM, and avoid the drug resistance of S. aureus (F Luo, Zeng, Yang, et al., 2021). The length of the non-polar terminal carbon chain and the number of unsaturated bonds of phospholipids used in MVL preparation may affect the tightness and fluidity of the liposome bilayer membrane, which would lead to the problems of sedimentation, aggregation, and drug leakage during MVL storage, thereby affecting the stability, entrapment efficiency, and drug release rate of MVL (Lu et al., 2021). We suggest using phospholipids with strong antioxidant capacity, such as dioleoyl lecithin, or coating a layer of sodium alginate on the liposome to improve the stability and entrapment efficiency of MVL. Moreover, in order to obtain an ideal drug release rate, some proper excipients can be adopted to modulate the drug release (e.g. using trioleic acid glyceride to slow down or trioctanoic acid glyceride to accelerate the drug release) (Y Li et al., 2020).

2.1.2. Polymer micelles

Polymer micelles have become ideal dosage forms for improving the bioavailability of flavonoids due to their good stability, biocompatibility, and solubilization (X Ma et al., 2019). In previous studies, our research group found that the polymer micelles prepared by electrostatic adsorption of negative-charged DHM with positive-charged natural polymers such as chitin and chitosan could improve the solubility and stability of the drug. Solutol®HS15 (HS 15) was a novel amphiphilic carrier material reported to have low toxicity and good solubilization (Nair et al., 2020). HS 15-loaded DHM self-assembled polymer micelle (DHM-MS) prepared by Ye et al. significantly improved the solubility, sustained release, and intestinal permeability of DHM and also showed better antioxidant and liver protection in drunk rats (Ye et al., 2021). However, DHM-MS only showed passive targeting delivery, thereby we suggest that HS 15 can be combined with ligands targeting liver tissue such as glycyrrhizic acid, or targeted by pH and inflammatory stimulation to improve the accumulation of DHM-MS in the liver tissue (Stecanella et al., 2021). It is noteworthy that, polymeric micelles are usually heterogeneous mixtures, even above the critical micelle concentration (CMC). In addition, polymeric micelles generally suffer from their structural instability and may easily disassemble to free polymers under dilute conditions or when exposed to environmental variations (e.g. pH and ionic strength), which may lead to low stability and certain toxic/side effects of the drug-loaded polymer micelles (S Lv et al., 2020). In previous studies, we found that micelles formed by mixed materials could make full advantage of each material, resulting in better properties such as higher stability and drug loading (Y Sun, Li, et al., 2019). Therefore, it is expected to increase the stability of the micelle system if other surfactants with the same hydrophilic chain as HS 15 were used to form mixed micelles (Y Sun, Li, et al., 2019).

2.1.3. Protein nanoparticles

Compared with other drug delivery systems, the protein nano-carrier delivery system has low antigenicity, good biocompatibility, and biodegradability (Lim et al., 2017). Nevertheless, protein nanoparticles have some disadvantages such as complex preparation process, poor stability, and difficulty in controlling side reactions, which need to pay attention to in practical use (Kianfar, 2021). Zein has strong hydrophobicity and good self-assembly properties, which
makes it ideal for controlling the release of insoluble drugs (Kianfar, 2021). Sun et al. prepared nanoparticles with zein as the carrier material and sodium tyrosine as the stabilizer, and encapsulated DHM by hydrophobic interaction, resulting in DHM-embedded zein-caseinate nanoparticles (DZP) with good dispersion and stability. DZP not only improved the adhesion and retention time of DHM in mouse gastrointestinal tract but also significantly increased the bioavailability of DHM in rats compared with DHM suspension (CC Sun et al., 2020). Among various protein nano-carrier materials reported, albumin has many advantages such as water solubility, biodegradability, biocompatibility, non-toxicity, non-immunogenicity characteristics, and particularly tumor targeting ability, thereby it is a very promising protein nano-carrier material to ensure the chemotherapy efficacy and safety of anti-tumor drugs (Kunde & Wairkar, 2022). Protein nano-delivery system is easy to modify the surface, using magnetic materials, pH-responsive materials, or connecting targeting ligands, to further enhance the targeting and curative effect of drug-loaded nanoparticles (Aziz et al., 2022).

### 2.1.4. Nanocapsules

With the emergence of multidrug-resistant strains, the use of polymer nanocapsule delivery systems loaded with active compounds is considered as a promising strategy to overcome biomembrane resistance (Deng et al., 2020; Nikezic et al., 2020). The pharmaceutical nanocapsule is a nanosized drug-loaded spherical particle with core-shell morphology, which has the advantages of increased solubility and stability, increased efficiency of encapsulation, the possibility of reducing adverse effects of drugs, and reduction of therapeutic doses (Frank et al., 2015). However, there are still many safety problems in nanocapsules, such as the surface charge that may have a toxic effect on cells and some toxic reagents involved in the preparation process, which have hindered their further application (Lima et al., 2022). Eudravit Rs100® polymer is a positively charged material with strong mucosal adhesion and can provide higher adhesion for positively charged bacteria (Dalcin et al., 2017). DHM nanocapsules (NC-DHM) prepared by Dalcin et al. using Eudravit Rs100® can significantly reduce the formation of Pseudomonas aeruginosa biomembrane in the urinary catheters. Compared with free DHM, NC-DHM showed prolonged drug effects and stronger antibacterial activity (Dalcin et al., 2019). It is noteworthy that Eudravit Rs100® polymer can induce the death of the PBMC cell line (i.e. the most sensitive human cell line for evaluating the toxicity of various compounds), and lead to decreased cell viability and increased DNA damage. However, the addition of DHM into Eudravit Rs100® nanocapsules could not only avoid cytotoxicity and genotoxicity, but also prevent the pyrolysis and photolysis degradation of DHM, and maintain the antioxidant effect of DHM (Wei et al., 2022). Considering the strong antioxidant activity of DHM itself, we suggest that DHM could be added as an excipient into the nanocapsules of Eudravit Rs100® to prevent cytotoxicity when delivering other drugs (Guo et al., 2016).

In summary, the DHM nano-delivery system usually has excellent controlled drug release, special tissue targeting, and strong modifiability, so it is easy to obtain high drug bioavailability and efficacy. It should not be ignored that the preparation of DHM nano-preparations involves many toxic reagents, expensive excipients, and complex processes, thus researchers need to continue to study low-toxic, economic reagents and materials, and simplified processes. In addition, the previous research on DHM nano-delivery systems mainly focused on the preparation process and drug quality investigation but often ignored the further study of their pharmacological activity and safety, which should be paid more attention to in the future.

### 2.2. Microemulsion preparation

Compared with the nano-drug delivery systems, the microemulsion is another widely studied drug delivery system, which is excellent in improving oral absorption and preventing drug oxidation (Garavand et al., 2021). It has been reported that DHM microemulsion systems include ordinary microemulsion preparation, Pickering emulsion, and self-emulsifying drug delivery system.

Solanki et al. found that DHM microemulsion prepared with Capmul MCM as oil phase (DHM/ME-C), Cremophor EL as surfactant, and T-transcutol as co-surfactant would change from water-in-oil to bicontinuous phase with the addition of water, which regulated the dissolution and oral bioavailability of DHM (Solanki et al., 2012). Our group has prepared DHM microemulsions with different polarity oil phases and emulsifiers with different HLB values, during which we found that the selection of proper emulsifier and oil phase in the microemulsion system is very important. We also found that the dissolution of DHM in the weakly polar organic phase is better, probably due to its abundant hydroxyl groups, thereby weakly polar organic solvents (such as Capmul MCM) are expected to be better for preparing DHM microemulsions. The problem to be solved in the clinical application of DHM microemulsion is the irritation caused by the addition of a large amount of oil phase, emulsifier, and co-surfactant. Therefore, developing safe and non-irritative microemulsion components would be an important task for future research. In addition, the pH-dependent solubility of some drugs in the microemulsion preparation has also been a problem that needs to be solved in the future (Egito et al., 2021).

Pickering emulsion is a kind of emulsion with ultrafine solid particles as the emulsifier. It has excellent coalescence stability, loading capacity, controlled release, biodegradability, and biocompatibility, which can make up for the shortcomings of traditional emulsions (XM Li et al., 2020; Mwangi et al., 2020). The natural fiber extract cellulose nanocrystalline (CNC) is a stable and effective solid particle emulsifier with good biocompatibility and biodegradability. Pickering emulsion stabilized by CNC can achieve increased loading and sustained release of DHM and can improve the stability of DHM (P Shen et al., 2022). Lysozyme (LY) is a multifunctional natural single-chain protein, which not only acts as an antibacterial agent but also has many beneficial effects such as...
regulating intestinal flora and reducing inflammation (H Liu, Wang, et al., 2022). In the research of Geng et al., DHM was adsorbed on the surface of LYZ to form a three-dimensional network structure to stabilize Pickering high internal phase emulsion (HIPE). Therefore, Pickering HIPE loaded with DHM has the valuable properties of antibacterial effect, lipid oxidation resistance, and lutein protection (Geng et al., 2022). It has been reported that some toxic and side effects such as allergy, nerve injury, and renal injury can be caused by some particle stabilizers in Pickering emulsion, which necessitated further safety studies of Pickering emulsion in the future (Zhang et al., 2020).

Self-emulsifying drug delivery system (SEDDS) is a homogeneous mixture of drugs, co-emulsifiers, hydrophilic emulsifiers, and oil phase, which can spontaneously form O/W emulsion or microemulsion under gastrointestinal peristalsis or mild stirring in ambient temperature, but it has some defects such as instability and large gastrointestinal irritation (Salawi, 2022). However, the solid self-emulsifying drug delivery system (S-SMEDDS) can well overcome these defects (C Sun et al., 2018). Wang et al. prepared DHM-loaded liquid self-microemulsion (DHM/L-SMEDDS) with octyl and decyl glycerate as the oil phase and polyglycerol-6 monooleate and polyglycerol polyricinoleate as the co-emulsifiers. Then DHM-loaded solid self-microemulsion (DHM/S-SMEDDS) was prepared by adsorbing DHM/L-SMEDDS onto the solid absorbent Aerosil 300. In vitro and in vivo experiments showed that DHM/S-SMEDDS improved the antioxidant activity and bioavailability of DHM, and showed good fluidity and stability under various storage conditions within 10 weeks (Wang et al., 2019). Theoretically, the increase in the specific surface area, the number of voids, and the depth of pores in the particle may lead to the difficulty of drug dissolution and the spontaneous aggregation of drug molecules (i.e. the ‘aging’ of crystal nucleus). Therefore, the above issues should be considered to prevent the aging phenomenon of S-SMEDDS during preparation (Kim et al., 2014).

2.3. Gel preparation

The reported gel preparations of DHM included emulsion gel and hydrogel. Emulsion gel is a kind of emulsion with a soft-solid structure, which endows liquid oil with the ability of controlled drug release. Pickering emulsion gel is stabilized by solid particles, which can be used as a controlled release carrier for bioactive compounds while maintaining gel properties (S Li et al., 2020; P Lv et al., 2020). However, due to the varied physicochemical properties of different solid particles, the Pickering emulsion gel may have flocculation and coalescence problems (S Li et al., 2020). It was reported that DHM could form a three-dimensional network with triglyceride, so DHM can be used as a particle stabilizer of Pickering emulsion gel. The prepared DHM-Pickering emulsion gel could not only effectively protect nutritious food in the oil phase, but also bring out a variety of health promotion functions of DHM (P Lv et al., 2020; S Geng, Jiang, et al., 2021). Similarly, Geng et al. prepared a Pickering emulsion gel with DHM and high-amylose corn starch (HCS), in which DHM formed supramolecular complexes (DHM/HCS) with some amylose molecules on the surface of HCS particles by non-covalent interaction. DHM/HCS stabilized Pickering emulsion gels by establishing a relationship of ‘molecular interaction-particle characteristics-emulsion gel properties’, which enhanced the gel structure and emulsifying ability of Pickering emulsion gels (S Geng, Liu, et al., 2021). Because DHM emulsion gel has the characteristics of high nutrition and high stability, it could be considered for functional food in the future.

Hydrogels are a major nutrient delivery system with the advantages of pH-responsive delivery and controlled release. Protein-polysaccharide hydrogels have excellent biocompatibility, and their gelation often occurs when the polymer concentration is low (Narayanaswamy & Torchilin, 2019). Wei et al. prepared hydrogel with ovotransferrin (OVT) fiber and xanthan gum (XG) as the materials. OVT fiber has good biocompatibility. XG is a common food additive. The hydrogel carrier assembled by electrostatic association has high gel strength and viscosity, which can significantly increase the drug loading of DHM in the hydrogel and release the drug slowly (Wei et al., 2020). Therefore, protein-polysaccharide hydrogel provides a new idea for the nutrition delivery system of natural drug extracts. Nevertheless, protein-polysaccharide hydrogels may suffer from poor stability (e.g. melting, aggregation, and condensation) and uncontrollable drug release rates under some extreme conditions (e.g. high temperature, strong acid and alkaline conditions), which is worthy of more attention in future research (K Liu et al., 2021; Manzoor et al., 2022).

It is worth mentioning that traditional hydrogels can be mixed with other dosage forms to produce new functions. Dalina et al. prepared hydrogels with ethyl cellulose (EC) as the matrix material, nipagin as the preservative, propylene glycol as the wetting agent, and Tween 80 as the solubilizer. DHM nanocapsules (NC-DHM as mentioned in '2.1.4 Nanocapsules') were added to the hydrogels. This NC-DHM hydrogel exhibits photoprotective activity against ultraviolet radiation-induced DNA damage and has good safety for the skin (Dalina et al., 2021), thereby we suggest that it could be developed for the cosmetics industry.

2.4. Crystal

Drug cocrystal is formed by hydrogen bond self-assembly between active pharmaceutical ingredients and appropriate cocrystal former. Drug cocrystal can modify the physicochemical properties of drugs without changing the covalent structure of drug molecules (Kavanagh et al., 2019). DHM is an ideal candidate for cocrystal due to its multiple hydrogen bond receptors and donors. Therefore, the preparation of highly soluble cocrystal can be an effective strategy to improve the physicochemical properties and biopharmaceutical properties of DHM (Morales et al., 2017).

Wang et al. obtained (+)DHM cocrystal by co-crystallizing DHM with the of theophylline. Compared with racemic DHM, (+)DHM metabolized more slowly in mice and showed better anti-inflammatory activity and bioavailability (C Wang, Xiong,
In order to prevent the DHM cocrystal from precipitating during the dissolution and maintain its supersaturation concentration, suitable precipitation inhibitors should be used. For example, when PVP K30 is used as the precipitation inhibitor, the stability of DHM-caffeine cocrystal and DHM-urea cocrystal in the medium can be enhanced, and the supersaturation concentration can be maintained, thus improving the oral bioavailability of DHM (C. Wang, Tong, et al., 2016). The effects of optical isomers are often different. For example, (−)gossypol has significant anticancer properties, while (+)gossypol has almost no such biological activities (H Liu, Zhang, et al., 2022). Since the pharmacological effects of (+)DHM and (−)DHM can be different, it is necessary to develop monoisomer drug preparations in the future to reduce ineffective substances for patients and avoid potential adverse/side effects (Tong et al., 2015).

Drug-drug cocrystal consists of two or more active drug components, and the ideal balance of physicochemical properties can be achieved through the non-covalent interaction among different drugs (Shinozaki et al., 2019). Both DHM and pentoxifylline (PTX) have the effects of treating vascular diseases and anti-cancer. They can form DHM-PTX cocrystal hydrate by hydrogen bonding. After forming cocrystal, DHM and PTX can be released synchronously and have a synergistic anti-cancer effect on HepG2 cells in vitro (L Liu et al., 2022). Similarly, Li et al. prepared berberine hydrochloride (BER) and DHM as BER-DHM cocrystal, which not only improved the thermal stability and solubility of DHM but also showed a synergistic anticancer effect on HT29 cancer cells in vitro (P Li et al., 2020).

The melting point should be strictly controlled in the process of developing drug cocrystals. A high melting point will lead to poor solubility, while a low melting point will decrease the stability of drugs. Nevertheless, there are still some problems in drug cocrystal, such as the instability in supersaturation state, how to choose cocrystal ligands, and unclear absorption mechanism in vivo, which warrant further study (Chaves et al., 2020).

2.5. Gastric floating preparation

DHM can exist stably in low temperatures and weak acidic environments (pH 6.0). When pH drops from 8.0 to 6.0, the absorption of DHM is significantly enhanced, suggesting that DHM is easier to be absorbed under acidic conditions (Shen et al., 2015). After oral administration of DHM in rats, the maximum concentration of DHM in the gastrointestinal tract is much higher than that in the heart, spleen, liver, lung, and kidney. DHM is a natural extract with a low titer, which often needs a large dose for administration to produce a curative effect. Based on the above facts, we suggest the gastric floating dosage forms may be ideal for DHM preparations (Tong et al., 2015).

After oral administration, the volume of gastric floating preparations can expand under the action of the gastric environment, making its density less than that of gastric contents. Therefore, they can float in gastric juice, thus prolonging the residence time of the drug in the stomach (H Liu, Wang, et al., 2021). In our previous study, DHM sustained-release gastric floating tablets (DHM-GFTs) were prepared by using NaHCO₃ as the foaming agent, ethyl cellulose (EC) as the adhesive, povidone K30 (PVP K30) and hydroxypropyl methylcellulose K4M (HPMC K4M) as the hydrophilic gel matrix. Compared with DHM powder, DHM-GFTs had a long-term gastric floating ability and sustained release for over 12 h, which significantly prolonged the residence time of DHM in rabbits and improved the oral bioavailability (H Liu et al., 2019). However, the drug loading content of DHM-GFTs was low (about 24%) and could not effectively avoid the large fluctuation of the blood drug concentration. Then, we further prepared DHM gastric floating pills (DHM-GFPs). With the optimal formulation, DHM-GFPs showed a much higher drug loading content (about 33%). Moreover, the gastric floating time and sustained release time were prolonged to over 24 h, which significantly reduced the fluctuation of the blood drug concentration and resulted in a better anti-inflammatory effect (H Liu, Gan, et al., 2021). Despite the advantages above, some gastric floating preparations may have incomplete absorption due to gastric emptying, gastric peristalsis, and gastric contents, thus reducing the oral bioavailability of the drug (Rajora & Nagpal, 2022). To overcome these problems, the development of gastric floating preparations with biological adhesion and mucinous adhesion, or using excipients with high density and high expansibility could be considered in the future (Iglesias et al., 2020).

2.6. Phospholipid complex

One of the reasons for the low oral bioavailability of drugs may be due to their limited solubility and/or low permeability (Kuche et al., 2019). Bombardelli et al. found that naturally extracted drugs have a special affinity with phospholipids, and they could form phospholipid complexes in water based on non-covalent interaction (Bombardelli, 1991). Phospholipid complex has amphiphilic characteristics, which can increase the solubility and stability of insoluble drugs, providing an effective method to solve the low oral bioavailability of naturally extracted drugs (Kuche et al., 2019). Liu et al. prepared a DHM-lecithin complex by solvent volatilization method. DHM and lecithin are completely combined by non-covalent bonds, producing a DHM-lecithin complex that could improve the solubility and antioxidant activity of DHM. As a natural antioxidant, DHM has the potential to replace butylated hydroxyanisole which is the most widely used synthetic antioxidant (Liu et al., 2012). Similarly, Zhao et al. prepared DHM phospholipid complex (DHM-HSPC com), in which DHM was dispersed in hydrogenated soybean phosphatidylcholine (HSPC) carrier in an amorphous state, so DHM-HSPC com significantly improved the solubility and drug loading content of DHM. In type II diabetes mellitus rats, DHM-HSPC com showed better hypoglycemic activity in vivo compared with the same dose of free DHM, and decreased the clearance rate and apparent distribution volume of DHM in vivo, thus improving the bioavailability of DHM (Zhao et al., 2019). Therefore, the DHM phospholipid complex would be a promising pharmaceutical preparation to improve the
bioavailability of DHM. When preparing phospholipid complex, the drug/lipid ratio would affect the drug binding rate, thereby choosing the appropriate drug/lipid ratio is the key factor to improving the drug loading in phospholipid complex (Biswa et al., 2019). Despite the great potential of the phospholipid complex technology, the instability (e.g. risk of aggregation and chemical degradation) and the drug leakage that can lead to irregular pharmacokinetics are still major issues which need further studies to settle in future studies (Kuche et al., 2019).

2.7. Cyclodextrin inclusion complex

Cyclodextrins (CD) are a kind of semi-natural cyclic oligosaccharides with conical hollow structures, which can improve the solubility of drugs. The whole B ring and part of the C ring of DHM could be embedded into the cavity of CD through non-covalent interactions such as hydrogen bonding (YP Wu et al., 2020). Therefore, inclusion technology may be an effective strategy to improve the low water solubility of DHM. Particularly, the hydroxypropyl-β-cyclodextrin (HP-β-CD) not only has the good properties of β-cyclodextrin (β-CD) such as low price and good safety but also has the characteristics of biodegradability and high solubility (Mu et al., 2022). Ruan et al. prepared DHM inclusion complexes DHM/HP-β-CD and DHM/β-CD using HP-β-CD and β-CD respectively, in which the solubility and dissolution of DHM were positively correlated with the addition amount of HP-β-CD and β-CD (Ruan et al., 2005). In the hyperlipidemia zebrafish model, DHM/HP-β-CD and DHM/β-CD showed better lipid-lowering activity than free DHM. In particular, DHM/HP-β-CD could induce the apoptosis of human HepG2 cells in a dose-dependent manner (Yang et al., 2011). It is noteworthy that although DHM/HP-β-CD showed good solubility and thermal stability, Liu et al. found that the antioxidant capacity of DHM was weakened due to the shield of HP-β-CD, which necessitated further research of DHM/HP-β-CD to settle this problem (Liu et al., 2012). Moreover, attention should also be paid to the problems of poor stability in acidic or alkaline environments, the presence of residual organic solvents, low drug encapsulation efficiency, and complex preparation process, which have limited the application of cyclodextrin inclusion complexes (Cid-Samamed et al., 2022).

2.8. Others

In addition to the above-mentioned dosage forms for DHM, there are still several reported DHM dosage forms such as solid dispersions, covalent polymers, and composite active packaging films. Solid dispersions are one of the most promising strategies to improve the oral bioavailability of poorly water-soluble drugs (Tran & Park, 2021). The solid dispersion prepared by Ruan et al. with hydrophilic polymers PVP K30 and PEG-6000 as carriers significantly improved the solubility and dissolution rate of DHM (Ruan et al., 2005). However, the stability of the solid dispersion and its easy absorbing water can be two problems that lead to phase separation, crystal growth, or crystal conversion during storage, which may further result in decreased solubility and dissolution rate (Cid et al., 2019). Liu et al. prepared DHM-sugar beet pectin (SBP) covalent polymer (SBP/DMH), in which DHM improved the emulsifying ability of SBP through non-covalent interaction, while SBP increased the solubility and antioxidant performance of DHM. Because SBP/DMH had a strong protein binding ability, it could be used as a noncompetitive inhibitor and anti-competitive mixed inhibitor of α-glucosidase, and its inhibitory effect is stronger than free DHM, and it exhibited the potential to treat non-insulin-dependent diabetes mellitus (Liu et al., 2018). Based on this, the covalent polymers formed by DHM and some natural polysaccharides can be potential ingredients to develop functional foods. Flavonoid-grafted polysaccharides can not only change the structural characteristics of polysaccharides but also improve the physicochemical properties and biological characteristics of polysaccharides. However, studies on the applications of flavonoid-grafted polysaccharides are very limited, and the safety of flavonoid-grafted polysaccharides is often ignored (J Liu et al., 2020; Bhanushali et al., 2022). Active packaging is able to inhibit microbial growth, enzymatic reaction, and oxidation reaction by adding various active substances into the packaging materials, so as to prolong the shelf life of packaged foods and improve their safety. In recent years, active packaging is considered as one of the best food preservation measures (Gulcin, 2020). Xie et al. prepared composite active packaging film (KGM/GG-DHM) with konjac glucomannan (KGM), gellan gum (GG), and DHM. KGM/ GG-DHM had good thermal stability, controlled release behavior, biocompatibility, antioxidant activity, and antibacterial activity against Escherichia coli and Staphylococcus aureus (Xie et al., 2021). The main shortcoming of active packaging is the uncontrollable migration rate of active compounds inside. Because of the low molecular weight of most active compounds, their release rates from active packaging are potentially rapid, in turn leading to a shorter shelf-life (Almasi et al., 2021). Therefore, it is still necessary to study the stability, release mechanism, and diffusion kinetics of active food packaging materials in the future, and further evaluate the timeliness of active components in the active packaging films.

In summary, the application of various techniques to prepare dosage forms for DHM can increase its solubility, permeability, and stability, which serves the needs of clinical treatment. This part introduced most of the reported DHM dosage forms, which were summarized in Table 1. In addition to the dosage forms introduced above, we can also consider the research of other dosage forms for DHM such as dropping pills, microcapsules, microspheres, etc., as well as the combination of various dosage forms in the future, so as to expand the application of DHM.

It is noteworthy that the current transformation of DHM research results into an application is not satisfactory. Although invention patents about DHM have been issued in various countries, so far only one DHM capsule has been approved for the market in the United States, which was probably due to the low titer of DHM that has led to
unsatisfactory drug effect when used alone (D Liu et al., 2019; J Shen et al., 2022). The combination of DHM and other drugs may be a promising strategy to expand the application of DHM. Apart from medicine, it should be noted that there are still very few reports on the application of DHM in the food industry. As mentioned in the previous parts, researchers can also consider combining DHM with a healthy lifestyle and developing it as the ingredient in functional drinks and foods.

3. Pharmaceutical designs for the combined application of DHM with other drugs

3.1. Combination with anti-tumor drugs

DHM has a variety of anti-tumor mechanisms. Researchers have found that DHM combined with other anti-tumor drugs can reduce multidrug resistance (MDR), toxicity, and side effects, and enhance the curative effects, therefore the

| Dosage forms | Structure diagrams | Labels | Main functions | References |
|--------------|--------------------|--------|----------------|------------|
| Liposomes    |                    | TS/CTS@DHM-lips | Enhancing the killing activity against E. coli and Staphylococcus aureus | (F Luo, Zeng, Chen, et al., 2021) |
|              |                    | Long-circulating DHM liposomes | Prolonging retention time, eliminating the half-life of DHM, and improving the bioavailability of the drug | (WJ Zhang et al., 2018) |
|              |                    | Multi-encapsulated DHM liposomes | Improving antibacterial activity and extending the antibacterial time | (F Luo, Zeng, Yang, et al., 2021) |
| Polymer micelles | DHM-Ms | Improving the gastrointestinal absorption and bioavailability of DHM, showing better antioxidant effect and protective effect on the stomach and liver against alcoholism | (Ye et al., 2021) |
| Protein nanoparticles | DZP | Improving the stability, diffusion rate, and bioavailability of DHM, as well as the adhesive property in mice gastrointestinal tract | (CC Sun et al., 2020) |
| Nanocapsules | DHM-loaded nanocapsules | NC-DHM | Sustaining drugs release and greatly reducing the amount of biofilm population in urinary catheters | (Dalcin et al., 2017) |
|              |                    |         | Maintaining the antioxidant capacity of DHM and preventing cytotoxicity or genotoxicity of the cationic polymer Eudralt Rs100 | (Dalcin et al., 2019) |

(Continued)
| Dosage forms                | Structure diagrams | Labels                        | Main functions                                                                 | References              |
|-----------------------------|--------------------|-------------------------------|-------------------------------------------------------------------------------|-------------------------|
| Microemulsion preparation   |                    | DHM/ME-C                      | Increasing the dissolution rate and oral bioavailability of DHM                | (Solanki et al., 2012) |
|                             |                    | DHM-CNC/Pickering Microemulsion | Improving the stability and sustained-release behavior of Pickering emulsion   | (P Shen et al., 2022)  |
|                             |                    | DHM-LY/HIPeS                  | Combining anti-bacterial, anti-lipid oxidation, and lutein protection          | (Geng et al., 2022)    |
|                             |                    | DHM/S-SED5                    | Improving stability, antioxidant activity, and bioaccessibility               | (Wang et al., 2019)    |
| Gel preparation             |                    | DHM Pickering emulsion gels   | Protecting oil-phase nutritional foods and giving Pickering emulsion gels new health-promoting properties | (Geng, Jiang, et al., 2021) |
|                             |                    | DHM/HCS-Pickering emulsion gels | Improving emulsifying capacity and stability                                  | (Geng, Liu, et al., 2021) |
|                             |                    | DHM/OVT fibril-XG hydrogel    | Improving drug loading, release efficiency, and oral bioavailability           | (Wei et al., 2020)     |
|                             |                    | N-DHM hydrogel                | As a photoprotective with antioxidant capacity                                | (Dalcin et al., 2021)  |

(Continued)
combination of DHM and other anti-tumor drugs have a good application prospect (Ferte et al., 1999).

Platinum drugs can combine with DNA to cause tumor cytotoxicity, which resulted in a high cure rate in clinical solid tumors (Zhang et al., 2022). However, due to their serious toxicity and side effects on normal tissues, especially kidneys, platinum drugs are limited in clinical application (Stankovic et al., 2020). It has been reported that DHM can be used as a reasonable chemosensitizer and attenuator of platinum drugs by activating the p53/Bcl-2 signal pathway, NF-kB/p65 signal pathway, and inhibiting the expression of multidrug resistance protein 2 (MRP2/ABCC2) (Figure 2) (Jiang et al., 2015; Wu et al., 2016; Wang et al., 2017). Adriamycin (ADR) is a highly effective anthracycline anti-tumor drug, which takes effects by inserting into the DNA of tumor cells. The clinical application of ADR is limited by disseminated intravascular coagulation (DIC) such as heart dysfunction and even heart failure (C Liu et al., 2020). Researchers have found that DHM can reverse MDR, reduce DIC, and enhance the curative effect by

| Dosage forms | Structure diagrams | Labels | Main functions | References |
|-------------|--------------------|--------|---------------|-----------|
| Crystal     |                    | DHM-caffeine cocrystal | Improving the physicochemical properties and biopharmaceutical properties of DHM | (C Wang, Tong, et al., 2016) |
|             |                    | (+)DHM-urea cocrystal | Improving anti-inflammatory activity and slowing down the metabolism of DHM | (C Wang, Xiong, et al., 2016) |
|             |                    | DHM-PTX·H2O | Improving solubility and synergistic anticancer effect on HepG2 cells in vitro | (L Liu et al., 2022) |
|             |                    | BER-DHM | Improving thermal stability and synergistic anticancer effects on HepG2 cells in vitro | (P Li et al., 2020) |
| Gastric floating preparation | | DHM-GFTs | Prolonging drug residence time in the body and improving bioavailability through good gastric floating ability and sustained drug release | (H Liu et al., 2019) |
|             |                    | DHM-GFPs | Extending the residence time of DHM in the body, improving bioavailability, and enhancing anti-inflammatory effect | (H Liu, Gan, et al., 2021) |
| Phospholipid complex | | DHM-lecithin complex | Improving lipid solubility, DPPH radical scavenging activity, and antioxidant activity | (Liu et al., 2012) |
|             |                    | DHM-HSPC com | Improving the C_{max} oral bioavailability while decreasing the clearance rate, apparent volume of distribution | (Zhao et al., 2019) |
| Cyclodextrin inclusion complex | | DHM/β-CD | Improving solubility and dissolution | (Ruan et al., 2005) |
|             |                    | DHM/HP-β-CD | Inhibiting HepG2 cell proliferation and inducing apoptotic cell death in a dose-dependent manner | (Yang et al., 2011) |
|             |                    | DHM/HP-β-CD | Improving solubility and thermal stability | (Liu et al., 2012) |
| Solid dispersions | | DHM/PVP-k30 | Improving solubility and dissolution rate of DHM | (Ruan et al., 2005) |
|             |                    | DHM/PEG 6000 | Improving solubility and dissolution rate of DHM | (Ruan et al., 2005) |
| Active films | | KGM/GG-DHM | Improving thermal stability, water resistance, UV-blocking ability, sustaining drug release behavior, and enhancing the antioxidant and antimicrobial activity of DHM | (W Xie et al., 2021) |
inhibiting activation of NLRP3 inflammatory corpuscle, P-gp function, MAPK/ERK signal pathway, p38MAPK signal pathway, and p53 signal pathway (Figures 3 and 4) (Zhao et al., 2014; Zhu et al., 2015; Xu et al., 2017; Y Sun et al., 2018; Y Sun, Liu, et al., 2019; Z Sun et al., 2020). Therefore, the combination with DHM is an effective way to solve the above problems of ADR. In addition to platinum drugs and ADR, DHM can also improve the efficacy of all-trans retinoic acid and reverse the MDR of 5-fluorouracil through various mechanisms (Zhou et al., 2014; He et al., 2018; Wu et al., 2020).

Although the combination of DHM and other anti-tumor drugs could lead to good coordination in mechanism, there are still some problems in the design of dosage forms, such as the differences in the physicochemical properties, absorption sites, pharmacokinetic behaviors, and effective doses of different drugs, resulting in the failure of the combination strategy (Santana et al., 2020). Therefore, when designing the combined preparations, we should consider the above problems and choose the appropriate ratio of drugs and dosage forms to achieve better synergistic drug efficacy. For example, the nano delivery system itself has a unique size effect and is easy to be functionalized, so it can enrich the drugs in tumor sites (Huang et al., 2022). Xie et al. have developed a nano delivery system for the co-delivery of curcumin (CuR) and methotrexate (MTX). When exposed to an aqueous solution, the DSPe-PeG-Imine-MTX pro-drug could spontaneously self-assemble into MTX-Imine-M nanoparticles encapsulating CUR within their hydrophobic DSPE core (Xie et al., 2018). The MTX-Imine-M-CUR nanoparticles could release the active form of MTX and CUR more efficiently in acidic media (pH 5) and exhibited significantly stronger tumor inhibition efficiency than the combination of both free drugs in vivo. Because the properties of nanoparticles in a water environment are more easily deteriorated than in a solid environment, solvent removal has become a potential strategy to improve the storage stability of drug-loaded nanoparticles (Franze et al., 2018). Previously, we prepared a solid mixture film (film-injection) consisting of the drug and the loading materials which could form drug-loaded nanoparticles when added with the proper solvent. The film-injection was prepared using the solvent evaporation technique and stored in a nitrogen atmosphere, minimizing the effect of the outside environment. This novel dosage form could significantly improve the long-term storage stability of nanomedicines and could be produced on a large scale (H Liu et al., 2020).

### 3.2. Combination with other drugs

In addition to anti-tumor activities, DHM also has many activities such as antiviral, antibacterial, liver protection, neuroprotection, antioxidant and anti-inflammatory, so it can also be used in combined therapy for other diseases (Wang et al., 2022). For example, the ortho-trihydroxy group in the B ring of DHM is the key pharmacophore to inhibit the activity of SARS-CoV-2 3CLpro which is an important target of coronavirus diseases, thereby the combination of DHM and SARS-CoV-2 3CLpro inhibitors such as acetooxyketone provides a new strategy to treat COVID-19 (Xiao et al., 2021; Xiong et al., 2021). In addition, DHM can inhibit ROS accumulation through pGC-1α/SIRT3 signal pathway and protect cells from apoptotic cell death, thus preventing hearing loss caused by oxidative damage of auditory hair cells (Han et al., 2020). DHM can also reverse the reduced bioavailability of ivermectin (IVA) due to the presence of P-glycoprotein, which is an innovative therapy for alcohol use disorder and secondary alcoholic liver disease (Silva et al., 2021). Moreover, the use
of DHM as an adjunct to BACE1 inhibitors in the treatment of Alzheimer’s disease (AD) can reduce the cross-inhibitory toxicity of BACE1 inhibitors, and can delay or reverse the AD progression (Das et al., 2020). In addition, DHM and ellagic acid may have a synergistic protective effect on UV-B damage of HaCaT cells and skin tissue by activating TGF-β1 and wnt signal pathways, which is the potential to protect photoaging (Moon et al., 2018).

In general, DHM in combination with multiple drugs may produce significant synergistic effects, as summarized in Table 2. However, the combined application of DHM was mostly for the treatment of cancer, while more studies for
other diseases should be considered. In addition, so far preparations for combined therapy of DHM with other drugs are scarcely reported, which necessitates the development of dosage forms for this application. When designing the dosage forms for combined therapies, not only the action mechanism but also the physicochemical properties and administration routes of the co-delivered drugs should be considered.

4. Some other issues to consider in the drug delivery of DHM

Due to the unstable structure of these natural products, the contact of DHM with metal ions, high temperature, and alkaline conditions should be minimized (Samsonowicz & Regulska, 2017). For example, the antioxidant effect of DHM depends on its phenolic hydroxyl groups in the B ring which
need protection during the production, processing, transporting, and storage. It is worth mentioning that most of the current studies are about the racemes of DHM. DHM contains two chiral carbon atoms, thereby theoretically there are four optical isomers. However, the pharmacological activity of the different optical isomers is yet to verify (Zuo et al., 2018). In order to avoid or reduce potential side effects from the non-drug parts, we suggest that the right single optical isomer DHM should be carefully selected for future research.

The therapeutic efficacy of traditional Chinese medicine is often the comprehensive effect under the simultaneous action of multiple components. The combination of various traditional Chinese medicines fully exhibited the unique advantages of multi-system, multi-channel, and multi-target treatment strategies (Y Xie et al., 2021). DHM has many pharmacological activities, but it alone has low titer and unsatisfactory effects, so it can be considered as an adjuvant to other drugs in the future, which provides a wider application prospect for this natural product. However, the current research on the combination of DHM and other drugs is basically blind and inefficient. To settle this issue, we suggest using network pharmacology and high-throughput screening to purposefully find and select appropriate combination strategies for DHM (Isgut et al., 2018). Based on this, the pharmaceutical design, feasibility, and safety of the combined application of DHM and other drugs should also be investigated.

5. Conclusion

Herein, the pharmaceutical design for DHM alone or with other drugs was reviewed. To date, various techniques have been adopted for producing all kinds of DHM preparations including nano-preparations, microemulsion, gels, crystal, and gastric floating preparation, etc., which has improved the solubility, permeability, and stability of this natural product. However, although invention patents about DHM preparations have been issued in several countries, the current transformation of DHM research results into marketable products is insufficient. The combined use of DHM with other drugs is a promising strategy to expand the application of DHM. So far, the combined applications are mostly for the treatment of cancer, while more studies are needed for the treatment of other diseases. For better synergistic effects, network pharmacology and high-throughput screening can be utilized to purposefully find and select appropriate combination strategies for DHM. Besides, preparations for combined therapy of DHM with other drugs are scarcely reported, which necessitates the development of dosage forms for this application. When designing the dosage forms for combined therapies, not only the action mechanism but also the physicochemical properties and administration routes of the co-delivered drugs should be considered. Apart from medicine, the development of DHM in the food industry is also of great potential. Due to its multiple effects and excellent safety, DHM preparations can be combined with a healthy lifestyle and developed as the ingredients in functional drinks and foods. Through this review, we hope to draw more attention from researchers in the related fields and provide valuable references for the research and development of other natural products with a similar structure-activity relationship to DHM. Although considerable time and costs are to consume in the future pharmaceutical studies of DHM, we believe that the research results of this natural compound will be gradually turned into market.
products with the development of new carrier materials and drug delivery systems.

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