High drug-loading nanomedicines: progress, current status, and prospects

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Abstract: Drug molecules transformed into nanoparticles or endowed with nanostructures with or without the aid of carrier materials are referred to as “nanomedicines” and can overcome some inherent drawbacks of free drugs, such as poor water solubility, high drug dosage, and short drug half-life in vivo. However, most of the existing nanomedicines possess the drawback of low drug-loading (generally less than 10%) associated with more carrier materials. For intravenous administration, the extensive use of carrier materials might cause systemic toxicity and impose an extra burden of degradation, metabolism, and excretion of the materials for patients. Therefore, on the premise of guaranteeing therapeutic effect and function, reducing or avoiding the use of carrier materials is a promising alternative approach to solve these problems. Recently, high drug-loading nanomedicines, which have a drug-loading content higher than 10%, are attracting increasing interest. According to the fabrication strategies of nanomedicines, high drug-loading nanomedicines are divided into four main classes: nanomedicines with inert porous material as carrier, nanomedicines with drug as part of carrier, carrier-free nanomedicines, and nanomedicines following niche and complex strategies. To date, most of the existing high drug-loading nanomedicines belong to the first class, and few research studies have focused on other classes. In this review, we investigate the research status of high drug-loading nanomedicines and discuss the features of their fabrication strategies and optimum proposal in detail. We also point out deficiencies and developing direction of high drug-loading nanomedicines. We envision that high drug-loading nanomedicines will occupy an important position in the field of drug-delivery systems, and hope that novel perspectives will be proposed for the development of high drug-loading nanomedicines.

Keywords: nanomedicines, nanocarriers, high drug loading, fabrication strategy, optimum proposal

Introduction

Conventional free-drug-treatment methods mainly involve oral, subcutaneous, and intravenous administrations. A drug spreads all over the body via blood circulation, and accumulates to a certain concentration in the target location to achieve a therapeutic effect. However, the desirable effects of drug treatment often decrease, due to the disadvantages of free drugs, such as the short half-life in the body, high dosage, poor selectivity, side effects on normal tissues, and multidrug resistance. Additionally, considerable amounts of marketed drugs and many new chemical drug candidates (as much as 70%) are neither absorbed through the intestines nor circulated by intravenous injection because of low water solubility. For instance, paclitaxel (Ptx), an effective anticancer drug, has a solubility of ~1 μg/mL in water, docetaxel, acknowledged as a drug with good solubility, has slightly higher solubility of 6–7 μg/mL in water. Therefore, solubilizing agents, such as dimethyl sulfoxide or surfactants, are typically...
added for systemic administration. However, such agents can still lead to neurotoxicity or other toxicities, even at very low dosages, thereby restricting the applications of free drugs.

The application of nanomedicines, such as inert carrier-based nanomedicines, polymer–drug conjugates (PDCs), coordination polymer nanoparticles (NPs), and carrier-free nanomedicines, can partly overcome the potential problems of free drugs in therapy. Excellent properties of nanomedicines are simultaneously or sequentially obtainable as follows: 1) expanded application ranges of hydrophobic drugs as a result of enhanced water solubility; 2) long circulation times of drugs when dosed into the circulatory system; 3) targeted delivery of drugs at the sites of the diseases through nonspecific targeting, such as the enhanced permeability and retention (EPR) effect or specific NP–cell surface interactions; 4) responses to local stimulation of the target location, such as abnormal pH, temperature, and ionic strength; 5) synergistic therapy via codelivery of multiple drugs at the same time and location; and 6) real-time monitoring of carrier (and drug) biodistribution and targeted accumulation.

Since the first US Food and Drug Administration (FDA) approval of nanomedicine (liposomal amphotericin B in 1990), more than 24 nanomedicines have been approved for clinical use, with total sales exceeding US$5.4 billion. The average cost and time required to develop a new nanomedicine (approximately $20–$50 million and 3–4 years) are significantly lower than for a new chemical drug (approximately $500 million and over 10 years). Therefore, the design and development of novel nanomedicines have higher market value and application prospects than that for conventional chemical drugs.

According to the fabrication strategies of nanomedicines, high drug-loading nanomedicines can be divided into four main classes (Scheme 1): 1) nanomedicines with inert materials as carriers, including inorganic porous carrier-based nanomedicines.
nanomedicines and absorption/desorption-type metal–organic framework (MOF)-based nanomedicines; 2) nanomedicines with drugs as part of the carrier, including linear and branched PDCs and infinite coordination polymer (ICP) I-type nanomedicines; 3) carrier-free nanomedicines, including drug nanocrystals (DNCs), amphiphilic drug–DCs (ADDCs), metal–biomolecule frameworks (MBioFs), and ICP II-type nanomedicines; and 4) nanomedicines following niche and complex strategies, such as aqueous noncovalent assembly and multiple assembly of drugs. For a visual comparison, the drug-loading content and drug-loading efficiency cited in this review are expressed according to the following equations:

\[
\text{Drug loading content (weight [wt]%) =} \frac{\text{Mass of the drug in nanomedicines}}{\text{Initial mass of the nanomedicines}} \times 100\% \tag{1}
\]

\[
\text{Drug loading efficiency (wt%) =} \frac{\text{Mass of the drug in nanomedicines}}{\text{Mass of the drug in feed}} \times 100\% \tag{2}
\]

Drug-loading content and drug-loading efficiency are two important parameters of nanomedicines. Drug-loading content reflect the mass ratio of drugs to nanomedicines, and drug-loading efficiency reflects the utilization of drugs in feed during the nanomedicine-preparation process. In essence, drug-loading content is determined by the structure and physical and chemical properties of the carrier material; drug-loading efficiency is determined by the drug-loading mechanism, mass of the drugs in feed, and other experimental conditions. It is more difficult to obtain high drug-loading content than high drug-loading efficiency for most nanomedicines. Through physical and electrostatic adsorption, the drug-loading process often results in low drug-loading efficiency, while through crystallization and covalent and coordinate bonds it often results in high drug-loading efficiency. Nanomedicines with high drug-loading content are usually prepared using processes with high drug-loading efficiency. If the space capacity of the carrier material is low, it is difficult to get high drug-loading content, even if the preparation process of nanomedicines shows high drug-loading efficiency. Taking into account most of the concepts investigated, high drug-loading efficiency may be necessary but not sufficient for a gain in high drug-loading content. In this paper, we focus mainly on drug-loading content, because this parameter is closely related to drug metabolism, side effects, extra burden, and therapeutic effect of the nanomedicines in vivo.

In most existing nanomedicines, inert carrier materials are mainly responsible for low drug-loading content (generally less than 10%). To administer a required drug dose in a target location, copious levels of carrier materials have to be used. For intravenous administration, the extensive use of these carrier materials may cause systemic toxicity, such as lipotoxicity, and impose an extra burden for patients to degrade, metabolize, and excrete these carriers. As such, FDA approval for most carrier-based nanomedicines is challenging to obtain, thereby limiting their clinical applications. On the premise of guaranteeing therapeutic effect and function, reducing or avoiding the use of carrier materials is a direct way of solving this problem. Moreover, it is easier for high drug-loading nanomedicines to achieve multidrug codelivery and combination therapy, which benefit from high drug loading. Given the advantages of combination therapy, including maximum therapeutic effect, minimum delay in induction of drug, and potential overcoming of multidrug-resistance mechanisms of tumors, multidrug-loaded high drug-loading nanomedicines will become a new hot spot in the field of drug-delivery systems. Although considerable efforts have been devoted to the fabrication of various high drug-loaded nanomedicines, the therapeutic effects are not always satisfactory and many theoretical and actual problems remain unsolved. In addition, not enough attention is paid to the study of high drug-loading nanomedicines, and innovative strategies are seldom reported. Therefore, the research status of high drug-loading nanomedicines must be reviewed and summarized, and their prospects should be discussed. In this review, the current research development of high drug-loading nanomedicines is investigated. The features of each fabrication strategy and proposals for possible optimization are discussed in depth. The inevitable role of high drug-loading nanomedicines in future drug-delivery systems is highlighted. We also hope that this study will provide us with a new perspective that will lead us to design and develop novel high drug-loading nanomedicines.

**Nanomedicines with porous material as carrier**

In this section, nanomedicines with porous materials, including inorganic porous materials, MOFs and protein NPs as carriers, are investigated. All these carrier materials provide high surface areas, large pore volumes, and feasible drug-loading methods, which contribute to high drug-loading capacity. The advantages and disadvantages of each material, as well as optimization options, are described.

**Inorganic porous materials as carriers**

To date, different kinds of inorganic porous materials, including mesoporous silica-based NPs (MSNPs), mesoporous
carbon NPs (MCNPs), magnetic colloidal NCs (MCNCs), mesoporous TiO₂ NPs,²⁵ and other inorganic porous materials, have been used to fabricate high drug-loading nanomedicines based on their intrinsic properties, such as large hollow interior, porous surface, and high surface:volume ratio. As summarized in Table 1, nanomedicines with inert material carriers can deliver different kinds of guest molecules.²⁶ In general, drug molecules are loaded into nanocarriers with high loading content through noncovalent electrostatic, π–π stacking, hydrogen bond, and hydrophobic interactions with carrier materials. In recent studies, drug molecules have been conjugated with the porous surface of carrier materials to acquire further increased drug-loading content and additional stimuli-responsive ability.

### Mesoporous silica-based materials as carriers
Since Vallet-Regí et al proposed MCM41 as a drug-delivery system in 2001,²⁷ MS-based materials, including MS, mesoporous calcium silicate, and mesoporous magnesium silicate, have been widely used in the fabrication of high drug-loading nanomedicines because of their intrinsic structure properties. The surface area and pore-volume ratio of MSNPs often exceeds 1,000 m²/g and 0.5–1 cm³/g, respectively, which facilitates the loading of large amounts of drug molecules.²⁸ Li et al showed that MSNPs could effectively load ibuprofen, a typical anti-inflammatory drug, with high drug-loading content of up to 21.6%.²⁹ Moreover, MSNPs present a site-selective controlled-release pattern by surface functionalization via any of the following three methods: cocondensation (one-pot synthesis), grafting (postsynthesis modification), and imprint coating.³⁰–³² For example, MSNPs functionalized with chitosan have a pH-triggered release characteristic, due to the swelling effect of chitosan in acidic environments.³³ This nanomedicine was used to deliver an oral sustained drug, carvedilol, with loading content and efficiency reaching 32.49±1.57% and 96.25±5.12%, respectively. Coincidentally, MSNPs exhibit temperature-responsive properties through a pore-capping copolymer poly-[N-isopropylacrylamide-co-1-butyl-3- vinyl imidazolium bromide].³⁴ When the temperature changes between 36°C and 40°C, the pore copolymers of MSNPs swelled or shrank to load or release the encapsulated cytochrome C. The drug-loading content of cytochrome C in this nanomedicine was 26.3% (Figure 1).

To enhance drug-loading content further, MSNPs with enlarged nanopores (E-MSNPs) and hollow MSNPs

### Table 1 Overview of two strategies of fabricating high drug-loading nanomedicines with porous materials as carriers

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<td>Inorganic porous materials as carriers</td>
<td>MSNPs/MCSNPs/MMSNPs</td>
<td>Noncovalent electrostatic and/or π–π stacking and/or hydrogen bond and/or hydrophobic interaction with carry materials</td>
<td>Hydrophobic drugs mostly; peptides, and gene drugs sometimes</td>
<td>≤69.3 wt%</td>
<td>11, 27–37, 39–42, 48</td>
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<tr>
<td>MOFs as carriers</td>
<td>MOFs</td>
<td>Coordinate bond and/or π–π stacking and/or hydrophobic interaction</td>
<td>Hydrophobic drugs mostly; gas drugs sometimes</td>
<td>≤58.3 wt%</td>
<td>108–120</td>
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<tr>
<td>Protein NPs as carriers</td>
<td>PNPs</td>
<td>Covalent bond with protein scaffold and/or π–π stacking, hydrophobic interaction with surface of PNPs and/or crystallization</td>
<td>Hydrophobically drugs mostly; peptides and gene drugs sometimes</td>
<td>≤44.9 wt%</td>
<td>122–126</td>
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Abbreviations: MCNPs, mesoporous carbon nanoparticles; MCSNPs, mesoporous calcium silicate nanoparticles; MMCNCs, mesoporous magnetic colloidal nanocrystal clusters; MMSNPs, mesoporous magnesium silicate nanoparticles; MOFs, metal–organic frameworks; MSNPs, mesoporous silica nanoparticles; PNPs, protein nanoparticles; wt, weight.

Figure 1 Preparation and temperature-dependent working mechanism of the cationic, thermosensitive, copolymer-capped MSNPs. Abbreviations: MSNPs, mesoporous silica nanoparticles; NIPAAm, N-isopropylacrylamide; BVIm, butyl vinylimidazolium.
(HMSNPs) can be selected as carriers. Compared with conventional MSNPs, E-MSNPs and HMSNPs have larger pore volume, which can accommodate more drug molecules.39 Eltohamy et al proved that E-MSNPs had higher drug-loading content for cytochrome C than normal MSNPs (50 wt% in E-MSNPs and 35 wt% in normal MSNPs, respectively).36 Similarly, Zhu et al57 proved that HMSNPs had significantly higher drug-loading content for aspirin than conventional MCM48 and MCM41, which were two nanocarriers discovered by Mobil researchers in 1992.34 The drug-loading contents of HMSNPs, MCM48, and MCM41 were 25.1 wt%, 19.2 wt%, and 16.6 wt%, respectively, under the same experimental conditions. In following study, they added polyelectrolyte multilayers to the surface of HMSNPs to fabricate a stimuli-responsive controlled-release system. The polyelectrolyte multilayers of sodium poly styrene sulfonate and polycationic poly(allylamine hydrochloride), prepared via a layer-by-layer method, yielded a drug-loading content of up to 41.64% for ibuprofen.39 Geng et al recently synthesized HMSNPs using hard template phenolic resin NPs to encapsulate two insoluble model drugs, namely carvedilol and fenofibrate.40 The resulting drug-loading contents for carvedilol and fenofibrate were 26.2±2.1% and 31.2±2.7% in normal MSNPs and 37.5±2.8% and 43.7±3.2% in HMSNPs, respectively. These findings indicate that HMSNPs have higher drug-loading content than normal MSNPs under the same conditions.

Another way to improve the drug-loading content of MSNPs is to conjugate drug molecules on the porous surface by covalent grafting or coordination bonds. Ebabe et al30 grafted two antioxidant molecules, caffeic acid and rutin, on to the surface of MSNP-NH$_2$R through amino groups on MSNPs and carboxylic groups in antioxidant molecules.41 They reported that caffeic acid had drug-loading content of 44.2%. Coordination is another feasible method. Farsangi et al prepared HMSNP-COOH carriers and loaded cisplatin in them via coordination bonds between carboxylic groups and Pt atoms in cisplatin molecules.42 These nanomedicines had a pH-dependent release pattern with drug-loading content as high as 48 wt%.

Mesoporous silicate-based NPs also have high drug-loading capacity. In the early research, calcium silicate is regarded as a candidate for bone replacement biomaterial due to its good bioactivity.43–46 Then, its mesoporous form is applicable in protein/drug delivery for bone regeneration because of its high surface area and large pore volume and good biocompatibility.47 Wu et al prepared ibuprofen-loaded mesoporous calcium silicate NPs with fine hierarchically nanostructure via a low-cost, surfactant-free sonochemical method.11 The drug-loading efficiency and drug-loading content of ibuprofen were as high as 86 wt% and 69.3 wt%, respectively. When the ibuprofen-loaded nanomedicine released the absorbed load, the calcium silicate material gradually transformed to hydroxyapatite, due to its good bioactivity. Similarly, Wang et al synthesized hollow mesoporous magnesium silicate NPs via a facile template-etching route.48 These well-prepared NPs exhibited a significantly high drug-loading content of 68.15 wt% and held a sustained-release property, showing great potential for cancer therapy.

Despite the encouraging progress made in the research of MS-based high drug-loading nanomedicines, several major challenges still need to be addressed, such as the in vivo toxicity caused by MS-based carrier materials. Although silica is considered generally recognized as safe and has secured FDA approval for Phase I human clinical trials,49 in vivo studies have described dose-dependent toxicity,50 such as inflammatory response,51 liver injury,52 neurotoxicity, and pregnancy complications.53 Studies have shown that particle size, surface charge,44 and surface functional groups also play significant roles in the cytotoxicity of MS-based carrier materials.54 Therefore, more careful and sufficient in vitro and in vivo studies are necessary to find the proper dose and type of MSNPs before clinical trials.

Mesoporous carbon as carrier

In contrast to the MSNPs already mentioned, MCNPs have even higher surface areas (as high as 2,000 m$^2$/g) and volume:mass ratios (>1 cm$^3$/g), which facilitate high levels of drug delivery.56–58 In early reports, MC microparticles were used to improve the oral absorption of water-insoluble drugs with drug-loading content higher than 50%.57,59–64 Applications of MCNP-based nanomedicines via intravenous administration have been quite limited, because of the following. First, ordered mesoporous carbons produced by carbonization of different chemical hydrocarbons usually have an undesirably size (>1 μm)$^{65,66}$ for drug delivery (<200 nm).67 Second, MCNPs exhibit poor dispersion in water under physiological conditions, due to their inherent hydrophobicity, which makes them impossible to use as carriers for intravenous drug administration. Finally, although some reports have shown that MCNPs have good biocompatibility,65,68 irregularly shaped MCNPs, such as carbon nanotubes or graphenes, can activate the complement system or induce immunotoxicity to some extent.99,100 With this limitation, MCNPs require further optimization to expand their application scope in drug-delivery systems. More recently, some MCNPs of <100 nm
diameter were obtained via template or hydrothermal method. Moreover, the hydrophilicity of MCNPs was improved by adding a hydrophilic cover, such as hyaluronic acid (HA) and polyethylene glycol (PEG), or introducing large amounts of oxygen-containing groups, such as hydroxyl, carboxyl, and sulfonic groups onto the surface. In one study, Liu et al successfully synthesized MCNPs with spherical morphology and uniform size (71 nm) via hydrothermal reaction of bacterial cellulose nanofibers (30–50 nm).12 These MCNPs disperse well in aqueous solutions, because of the introduction of a large amount of hydrophilic functional groups, such as –SO\(_3\)H, phenolic –OH, and –COOH groups, onto the surfaces. The drug-loading efficiency and drug-loading content of this nanomedicine for doxorubicin (Dox) were 93.4 wt% and 52.3 wt%, respectively. In another work, spherical oxidized MCNPs with an average diameter of 90 nm were prepared through low-concentration hydrothermal treatment and mild oxidization. Then, PEGylated phospholipid 1,2-distearoyl-sn-glycero-3-phosphoethanolamine (DSPE)-methoxy-PEG\(_{2,000}\) was used as a protective layer added to the outer surface after Dox loading. The PEG layer on the surface prevented the premature and burst release of drugs in the circulation system, as well as helping to avoid recognition of the reticuloendothelial system.71,72 The drug-loading content of this nanomedicine for Dox was as high as 59.7%±2.6% under optimum conditions (Figure 2).73 Mesoporous carbon-based nanomedicines have two inherent advantages, due to their unique mesoporous structure and carbonaceous composition. First, mesoporous carbon-based high drug-loading nanomedicines often exhibit good pH sensitivity in drug-delivery systems. This feature is due to the fact that the adsorption of most anticancer drugs with aromatic structures, such as Dox, by carbonaceous structures via π–π stacking and hydrophobic interactions is pH-dependent. Second, MCNPs serve as efficient photosensitizers with high absorption in the near-infrared region.74–77 Therefore, they function as promising nanocarriers in photothermal and chemical combination therapy. Zhou et al78 used these advantages to develop an effective dual-triggered (Hyal1/redox) chemophotothermal synergistic therapy nanomedicine, which presented efficient heat transformation for photothermal therapy. Research on MCNPs in drug-delivery systems is still in its infancy, and MCNPs face the following unsolved problems: 1) scalable synthetic strategies and standard methodologies to obtain MCNPs with suitable size and hydrophilic surfaces for intravenous administration; 2) unsuitability of traditional oxidation methods in the surface modification of MCNPs, because such techniques partially destroy the carbonaceous framework and the mesostructure of MCNPs; and 3) the limited application scope of MCNPs in nanomedicines and absence of applications of MCNPs in the field of gene and protein drug delivery, as opposed to MSNPs.

Figure 2 Synthesis, drug loading, and surface modification of MCNPs.

Abbreviations: MCNPs, mesoporous carbon nanoparticles; DSPE, 1,2-distearoyl-sn-glycero-3-phosphoethanolamine; mPEG, methoxy polyethylene glycol; oMCNs, oxidized mesoporous carbon nanospheres; Dox, doxorubicin; PEG, polyethylene glycol.
Mesoporous magnetic colloidal nanocrystal clusters
Mesoporous/hollow MCNCs (MMCNCs/HMMCNCs) have drawn high attention in targeting drug-delivery systems during the past decade. HMMCNCs and MMCNCs are promising candidates for high drug-loading nanomedicines, because of their high magnetization, large surface area and pore volume, and excellent colloidal stability. Compared with other inorganic porous materials, MMCNCs and MMCNCs are known as the “magnetic motor” for site-specific drug-delivery applications.

The most common fabrication method of MMCNC-based nanomedicines begins with a combination of solvothermal reaction and electrostatic stabilization, and follows with drug loading via nanoprecipitation. Luo et al synthesized Ptx-loaded Fe$_3$O$_4$ MMCNCs using poly(γ-glutamic acid) as an electrostatic stabilization agent. The resulting MMCNC-based nanomedicine possessed high magnetization, large surface area (136 m$^2$/g) and pore volume (0.57 cm$^3$/g), excellent colloidal stability, prominent biocompatibility, acid degradability, and high drug-loading content of 35 wt% for Ptx.

The fabrication of HMMCNCs involves both template-based and template-free solvothermal reaction methods. Li et al synthesized an HMCNC-based Dox-loaded nanomedicine with porous shell and tunable hollow chamber by a one-pot solvothermal process. This HMCNC carrier had a high surface area of 41.2 m$^2$/g and pore volume of 0.36 cm$^3$/g, which contributed to high drug-loading capacity. Furthermore, receptor-specific pH-sensitive folate-modified polyacrylic acid (PAA-FA) was grafted on the surface of HMCNCs via a versatile ligand-exchange method, because folate molecules can target cancer cells that overexpress folate receptors and enhance the endocytosis capability of such cancer cells. This nanomedicine is stable under high pH and quickly releases Dox under low pH. This site-specific targeting delivery was achieved by protonation and collapse of the PAA layers after cell internalization. The NPs presented drug-loading content of 24 wt% for Dox (Figure 3).

As described, novel inert magnetic NCs are promising nanocarriers, and their drug-delivery application is potentially feasible. Exciting progress has been made in MMCNC cluster-based nanomedicines, because of their high magnetization and large surface area. However, the fabrication of magnetic nanocarriers that demonstrate enhanced biocompatibility and excellent colloidal stability remains a challenge, which is critical for the application of magnetically driven drug delivery. Currently, in vivo toxicity studies and pharmacological experiments are scarce, and further studies are necessary.

Absorption/desorption-type MOFs as carriers
MOFs, a class of coordination polymers, are another type of porous material (the other class of coordination polymer is an ICP particle that is covered in later sections). MOFs consist of transition metal ions/clusters and multidentate organic ligands, and are typically synthesized via coordination-directed self-assembly processes under mild conditions (Figure 4). Over the past two decades, MOFs have been applied in photonics, catalysis, biosensors, gas storage, and drug delivery. For drug delivery, the high drug-loading capacity of MOFs relies largely on three factors: 1) high surface area (3,100–5,900 m$^2$/g) and large, regular, accessible cages and tunnels; 2) good physicochemical stability in circulation and quick stimuli

**Figure 3** Dox encapsulation in HMCNCs and pH-stimulated release of Dox from folate–HMCNC-Dox.

**Abbreviations:** Dox, doxorubicin; HMCNC, hollow magnetic colloidal nanocrystal; PAA, polyacrylic acid.
response, especially pH response; and 3) selectable metal ions with low toxicity (ie, Fe or Zn) or metal ions associated with the mechanism of drug action (ie, Cu with gossypol).  

Two approaches can be considered in fabricating high MOF-based drug-loading nanomedicines. For absorption/desorption-type MOF-based high drug-loading nanomedicines, drug molecules are absorbed in cages and tunnels of MOFs. For MBioFs, the bioactive drug molecules function as ligands that coordinate with the central metal atom. In this section, absorption/desorption-type MOF-based nanomedicines are summarized, whereas the latter are outlined in a following section.

Férey et al first reported absorption/desorption-type MOF-based nanomedicines in 2004. A model drug, ibuprofen, was absorbed by two dehydrated MOFs – chromium-based MIL-100 and MIL101 effectively in hexane solution. X-ray powder diffraction and N$_2$-adsorption experiments showed that both MOFs retained their excellent crystalline structures, and almost all the cages and tunnels were filled and/or blocked by the drug molecules. The drug-loading contents of ibuprofen for MIL100 and MIL101 were 58.3 wt% and 2.6 wt%, respectively. The different absorption capacities between MIL100 and MIL101 were attributed to their fine structure inside the particles. The larger pore size of cages in MIL101 made it easier to absorb ibuprofen compared with MIL100. The size of ibuprofen is 6×10.3 Å, whereas some cages in MIL100 were 4.8×5.8 Å (pentagonal), so the latter were not accessible for ibuprofen. The adsorption mechanism of ibuprofen in MIL100 and MIL101 was investigated by Babarao and Jiang through computational studies. They suggested that a part of the ibuprofen molecule penetrates MIL101 by forming coordinate bonds with unsaturated chromium molecules in the surrounding regions, and additional ibuprofen molecules weakly interact (ie, π–π interactions and hydrophobic interaction) with existing ibuprofen molecules. The results of molecular simulation in this study were consistent with “two-stage” release profiles observed in Férey et al’s work. Similarly, small-molecule drugs or gas drugs can be either chemisorbed by forming coordination bonds with transition metals or physically adsorbed into coordination polymers by weak intermolecular forces in absorption/desorption-type MOF-based nanomedicines.

For biological application, MOFs can be modified on the surface to increase size, decrease aggregation, and exhibit targeting performance. Horcajada et al developed a series of iron(III)-based MOFs (MIL53, MIL88A, MIL88Bt, MIL89, MIL100, and MIL101$_{\text{NH}_2}$) with engineered cores and surfaces for retroviral drug delivery. As expected, most of these MOFs (ie, azidothymidine triphosphate-loaded MIL101$_{\text{NH}_2}$, cidofovir-loaded MIL101$_{\text{NH}_2}$, urea-loaded MIL100, and urea-loaded MIL53) displayed high drug-loading capacity of over 40 wt%. PEG chains with only one terminal reactive group (amino or carboxyl) were firmly bound to the nanomedicine surface through coordinate covalent bonds of its amino or carboxyl end group with the metal centers. The PEG coating might control the nanomedicine size, protect the nanomedicines from aggregation, and escape sequestration by the reticuloendothelial organs.

Despite the research progress made, the potential application of MOFs in drug delivery is still in its infancy compared with other nanocarriers. Three major existing problems impede MOFs as mature high drug-loading nanomedicine carriers: 1) the relationship between the pore size of cages
in different kinds of MOFs and the size of cargo drug molecules needs to be systematically studied and determined; 2) appropriate ligands and low-toxicity metal ions are needed to improve biocompatibility, and more efficient surface functionalization is needed to ensure longer blood circulation and better targeting effects; and 3) more careful and sufficient in vitro and in vivo experiments need to be conducted to expand the application prospects of MOFs in the biomedical field.

Protein NPs as carriers
As potential nanocarriers, protein NPs have intrinsic advantages, including synthesis in natural sources, precise size and uniformity, hollow inter structure, self-assembling ability, biocompatibility, and biodegradation. Hydrophobic drug molecules, peptides, and gene drugs are covalently conjugated to the hollow cavity of the protein scaffold or noncovalently loaded into the internal space. Generally, restricted by the number of attachment sites, most of the protein NP-based nanomedicines prepared by conventional chemical conjugation present the same drawback of lower drug-loading content (<10%). One feasible way to enhance the drug-loading capacity of protein nanocarriers is designing and increasing the hydrophobic surface area of the protein NP. Inspired by transport mechanism of multidrug-efflux transporters, Ren et al implemented a biomimetic approach to enhance hydrophobic drug–protein interactions. Several virus-like protein NPs were redesigned and prepared via self-assembly of the E1 component of the pyruvate dehydrogenase multienzyme complex. Phenylalanine, one of the most hydrophobic amino acids, was systematically introduced to increase the hydrophobicity of E1’s hollow cavity surface. The drug-loading content of protein NPs for Dox reached 13.4%, and the theoretical maximum loading content was thought to be as much as 44.9%.

Although using protein NPs in the fabrication of high drug-loading nanomedicines is potentially feasible, a major concern with protein nanocarrier-based nanomedicines is their immunogenicity. When these nanomedicines are administered for the second time, elevated immune responses can occur and induce rapid clearance or neutralization or create potentially severe inflammatory responses. One way to bypass this problem is surface functionalization by introducing PEG and other coatings to reduce immunorecognition. The other is harnessing this natural effect and using it in immunobased treatments.

Nanomedicines with drugs as part of carrier
Although absorption/desorption-type high drug-loading nanomedicines with inert porous carriers have attracted heightened interest in fabricating high drug-loading nanomedicines, the inherent limitations of this strategy cannot be ignored. Drug loading and release are mostly achieved by absorption and desorption, which strongly depend on the diffusion rate of drug molecules and drug–carrier interactions in the physiological medium. In the stage of nanomedicine design, the interactions are difficult to predict. Also, the degradation of the inert carrier material may lead to additional toxicity and impose an extra burden on patients to excrete the carriers.

For drugs with stable chemical properties in the circulatory system, the protective function from carriers is not essential. To overcome this limitation and further improve the drug-loading content of nanomedicines, the direct way is gradually reducing or replacing the use of carrier material. With this goal in mind, two popular strategies are proposed to fabricate such high drug-loading nanomedicines. One strategy is to form prodrugs by conjugating drug molecules with polymers, and another is to form ICP I-type nanomedicines, in which drug molecules function as ligands and coordinate with the central metal atom (Table 2). The drug and polymer are two indispensable excipients in high drug-loading nanomedicine. Without one of these, the nanostructures cannot be maintained.

Table 2 Overview of two strategies of fabricating high drug-loading nanomedicines with drug as part of carrier

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<td>Branched polymer–drug conjugate</td>
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<td>ICP I-type nanomedicines</td>
<td>ICP I-type nanomedicines</td>
<td>Coordinate bond</td>
<td>Drugs with complexing ability (both hydrophobic drugs and hydrophilic drugs)</td>
<td>Not shown</td>
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Abbreviations: ICP, infinite coordination polymer; wt, weight.
Polymer–drug conjugates

Compared with the physical encapsulation of drug molecules in inert carriers, PDCs, such as linear and branch types, are excellent candidates for high drug-loading nanomedicines, because of their limited usage of inert carrier materials and self-assembly abilities. In addition to the common advantages of other nanomedicines already mentioned, the smart covalent links between the drugs and polymers endow them with multiple advantages: 1) stimuli-triggered release and targeted delivery; 2) protection of drugs from leakage and premature release during intravenous administration; 3) altered drug pharmacokinetics in the whole organism and even at the subcellular level, which may enhance the drug’s therapeutic value; and 4) additionally loading hydrophilic drugs into the core of micelles to achieve multidrug-combination therapy. For stimuli-responsive PDCs, pH-responsive types are the most frequently used. Several pH-responsive covalent links, such as acetal, orthoester, hydrazone, imine, cis-acotinyl, and oxime, are common, due to their rapid hydrolysis in the endosomal compartment (pH about 5). Moreover, a reduction-response covalent link disulfide depends on different redox potentials between the extra- and intracellular environments, especially in tumor cells.128–130

Linear polymer–drug conjugates

As the name suggests, linear PDCs are block copolymers that have a simple linear monomeric structure. Shen et al formed dual-drug-loading nanomedicines by conjugating one or two strong hydrophobic camptothecin (CPT) molecules to a very short oligomer chain of oligoethylene glycol (OEG) to form biodegradable ester bonds (Figure 5).131 The amphiphilic prodrugs OEG-CPT and OEG-DiCPT self-assemble into 100–200 nm liposome-like nanocapsules in aqueous solutions. The drug-loading contents for CPT were 40 wt% and 58 wt% in OEG-CPT and OEG-DiCPT, respectively, with undetected burst releases for both. Meanwhile, the liposome-like structures further loaded a water-soluble Dox HCl salt via a dialysis mechanism similar to drug loading into conventional liposomes. The drug-loading efficiency and drug loading content of Dox HCl were 85% and 18.5 wt%, respectively. Once inside the cells, the Dox HCl-loaded amphiphilic CPT prodrug liposomes released both CPT and Dox HCl, thereby providing synergistic cytotoxicity to cancer cells, as confirmed by high in vitro and in vivo antitumor activities. Similarly, Zhang et al synthesized a nanomedicine by conjugating the hydrophobic drug SN38 (7-ethyl-10-hydroxy-CPT), the active metabolite of CPT, to a low-molecular-weight OEG via an ester bond. The amphiphilic OEG-SN38 monomer self-assembled into micelles with a hydrodynamic diameter of 28.74 ± 2.51 nm, and a high drug-loading content of 36 wt% for SN38 was obtained.132 Gou et al133 anchored hydrophobic Dox to a low-molecular-weight PEG chain via a pH-responsive covalent hydrazine bond to form a linear amphiphilic PDC – PEG-Dox. This amphiphilic prodrug self-assembled into micelles with a diameter of about 125 nm in aqueous solution, and the drug-loading content for Dox was as high as 46 wt%. Once the nanomedicines were taken up by cells via endocytosis, the hydrazine bonds were hydrolyzed in the acidic environment of the endosomal compartment, and the Dox molecules were simultaneously released.

Figure 5 Amphiphilic CPT–polymer conjugates (OEG-CPT and OEG-DiCPT) and their self-assembly into nanocapsules to load other hydrophilic drug.

Note: OEG-DiCPT means two CPT molecules were conjugated to one OEG molecule.

Abbreviations: CPT, camptothecin; OEG, oligoethylene glycol.
Furthermore, PDC prodrug micelles can achieve targeted delivery by introducing targeting ligands, such as carbohydrates, folate, transferrin, peptides, and HA, onto the surfaces. For example, Guo et al synthesized a linear PDC – FA-PEG – poly(ε-caprolactone (PCL) – hydrazone-Dox – to form pH-sensitive folate-functionalized nanomedicines. The FA-PEG-b-PCL-hydrazone-Dox conjugates self-assembled into micelles with a hydration diameter of 70.9 nm, according to dynamic light-scattering measurements. In these micelles, Dox molecules anchored to PCL via hydrazone bonds functioning as the hydrophobic part, whereas PEG served as the hydrophilic part. The FA layer on the surface of micelles facilitated the endocytosis of the folate receptor-overexpressing tumor cells.

**Branched polymer–drug conjugates**

Compared with linear PDCs, branched PDCs, including branched star PDCs, hyperbranched PDCs, dendrimer PDCs, and brush PDCs, exhibit controllable supramolecular morphologies, self-assembly behavior, and thermal, mechanical, rheological, and solution properties. All these properties render branched PDCs viable for high drug-loading nanomedicines.

Branched star polymers are branched polymers containing several linear chains attached to a central core. Star PDCs often have a compact structure, globular shape, and large surface; they also possess enhanced solubility, low melting viscosity, unique rheological properties, and a large number of drug-binding sites. All these excellent properties depend on the arm’s molecular weight. Kowalczyk et al synthesized core–shell–type star polymer–cisplatin conjugates with a branched poly(p-[iodomethyl]styrene) acting as a hydrophilic core and PAA arms functioning as a hydrophilic shell. These star-type PDCs exhibited “unimolecular micelle” behavior in aqueous solution, and were globular with a hydration diameter of 12.9–14 nm. The cisplatin molecules were loaded into the unimolecular micelles via the coordinative bonds between the platinum atoms of cisplatin molecules and high density of carboxyl groups on PAA arms. The drug-loading content for cisplatin was 45 wt%. In vitro experiments showed lower cytotoxicity that was consistent with the sustained release of cisplatin molecules and slower cellular uptake of conjugated prodrug micelles compared with free cisplatin. Some examples of star PDCs utilized for nanomedicine fabrication following this strategy have been presented in the literature.

In a recent study, Liang et al designed pH-sensitive star polymer–Dox conjugates, in which a triblock copolymer 2-(N,N-dimethylaminoethyl)-methacrylate-co-p-(methacryloxyethoxy)benzaldehyde-co-2,2-dithiodiethyl dimethacrylate served as a hydrophilic core and poly(6-O-methacryloyl-d-galactopyranose) arms acted as a hydrophilic shell. Instead of loading drug molecules on the arm as previously mentioned, Dox molecules in this nanomedicine were loaded into the hydrophilic core through acidic–labile imine bonds between the aldehyde groups in p-(methacryloxyethoxy)benzaldehyde and the primary amines in Dox molecules. The drug-loading content of Dox in this nanomedicine was controllable in the range of 7.3–37.6 wt%, depending on the different grafting degrees of Dox. These star polymer–Dox conjugates self-assembled into unimolecular micelles with average diameters of 30–50 nm in aqueous solution. Dox molecules were quickly released at pH 5 and 6 due to the acid lability of aromatic imine linkage between the Dox molecule and star polymer. All these properties were desirable for drug delivery.

Hyperbranched polymers (HBPs) are another kind of branched polymer with a highly three-dimensional dendrite-like architecture, and their degree of branching is 0.4–0.6 (0 for linear polymer and 1 for dendrimer). Similar to branched star polymers, HBPs have serious adjustable physicochemical properties, such as molecular entanglement, solution viscosity, solubility, host–guest interaction capacity, and self-assembly behavior, caused by their modifying branches, end groups, and controllable degree of branching. Given these beneficial properties, HBPDCs are excellent candidates for high drug-loading nanomedicine fabrication.

Prabaharan et al synthesized the first HBPD C nanomedicines in 2009. In their work, a folate-targeting unimolecular micelle nanomedicine was synthesized, namely Boltorn H40-poly(l-aspartate-doxorubicin)-b-PEG/FA-conjugated PEG (H40-P[LA-Dox]-b-PEG-OH/FA). This nanomedicine contained a commercial aliphatic dendritic polyester Boltorn H40 core, hydrophobic poly(l-aspartate) inner arm, and a hydrophilic PEG and FA-conjugated PEG outer arm. Dox molecules were conjugated to the inner arm through a pH-sensitive hydrazone linker, resulting in a drug-loading content of 16 wt% for Dox. This block polymer self-assembled into micelles with sizes of 17–36 nm under dynamic light scattering and 10–20 nm under transmission electron microscopy. Following this strategy, Ye et al fabricated HBP–cisplatin conjugates based on hyperbranched polyglycerols (HPGs).

In this system, HPGs were modified with hydrophobic alkyl (C8-C10) chains into the core, derivatized with methoxy PEG (MePEG) onto the surface, and further functionalized with carboxylate groups on MePEG chains to conjugate and release cisplatin. The drug-loading efficiency of cisplatin was...
100%, and its drug-loading content was adjusted depending on the conjugated carboxylate levels. Respectively, the drug-loading content of HPG-C8/C10-MePEG6.5-COOH was 10 wt%, whereas that for HPG-C8/10-MePEG6.5-COOH348 was 20 wt%. These HBP-cisplatin conjugates self-assembled into micelles with hydrodynamic diameters of 5–10 nm. These conjugates demonstrated pH-triggered release behavior caused by the coordinate bond between cisplatin and carboxylate groups on MePEG chains. In vitro experiments showed that these nanomedicines presented good biocompatibility and antitumor effects.

Segmented polymers are a large family. Depending on the chemical differences among main chains, side chains, and grafting density of the side chains, the family of segmented polymers is classified as follows: graft polymer, in which the side chains have a different chemical nature from the main chains and low grafting density; brush-like polymers, in which the side chains have a different chemical nature from the main chains and high grafting density; and comb-like copolymers, in which the side chains have a similar chemical nature to the main chains and lower grafting density than brush-like polymers. Graft and brush-like polymers are excellent candidates for high drug-loading nanomedicine fabrication, depending on their side chains’ selectability and modifiability. Xu et al reported a novel graft polymer–Dox conjugate with a Dox-loading content as high as 40 wt%, namely, MPEG monomethylether-b-poly(methacrylamide tert-butyl carbazate-Dox) (MPEG-b-Dox) (Figure 6).

The MPEG in this conjugate functioned as the hydrophilic part, and methacrylamide tert-butyl carbazate, which was conjugated to Dox via an acid-labile hydrazine bond, acted as the hydrophobic part. The amphiphilic conjugates self-assembled into micelles with average hydrodynamic size of 80 nm. These conjugates also showed minimal drug release at pH 7.4 and high Dox release at pH 5, following the previously mentioned mechanism. In parallel, Zou et al synthesized a brush PDC, PLA-g-Ptx/PEG, through azide–alkyne click reaction of acetylene-functionalized PLA with azide-functionalized Ptx and PEG. The drug-loading content of Ptx reached nearly 23.2 wt%.

Similarly to linear PDC liposomes, branched PDC unimolecular micelles with hydrophobic inner domain can further effectively encapsulate hydrophobic substances into their cores for imaging or codelivery. Tai et al reported γ-CPT and Dox dual-drug-loaded nanomedicines prepared via polymerization of γ-CPT–glutamate N-carboxyanhydride on a PEG-based backbone via ring-opening polymerization. The amphiphilic conjugates self-assembled into unimolecular micelles with average hydrodynamic size of 50 nm. Dox molecules were then loaded into the hydrophobic core through physical interactions, such as π–π stacking. The drug-loading contents of CPT and Dox were 25.1 wt% and 30 wt%, respectively. An in vivo study demonstrated that dual-drug-loaded nanomedicines displayed significant synergistic effect on lung cancer-xenograft mice.

PDCs are an emerging one-component strategy for high drug-loading nanomedicine fabrication. This one-component feature with drug as part of carrier is essentially a double-edged sword: 1) for a particular PDC, the influence of the covalent links between drug molecules and biodegradable polymers to drug pharmacokinetics cannot be predicted; 2) the drug-release process can be complicated, which often involves both the nanostructure dissociating into monomeric units and the chemical breakdown of the linker; and 3) the choice of degradable linkers significantly affects the release mechanism of free bioactive drug from assembled nanostructures.

Figure 6 Formation of mPEG-b-Dox micelles.

Abbreviations: mPEG, methoxy polyethylene glycol; Dox, doxorubicin.
ICP I-type nanomedicines

ICPs are one of the classes of coordination polymers. Compared with conventional MOF-based nanomedicines, ICP-type high drug-loading nanomedicines often exhibit certain additional advantages. The ICP-type nanomedicines have a trimmable size and well-defined shapes,\(^\text{159}\) therefore, a desired particle size can be obtained for intravenous injection compared with absorption/desorption-type MOF-based nanomedicines and MBioFs. The pH-responsive point of nanomedicines can move to a targeted location without any additional dramatic alterations in the polymeric structure, due to the controllable degree of polymerization, chain monodispersion, intermolecular forces, and drug–metal coordination bonding strength.\(^\text{160,161}\) Several conventional anticancer drugs with autofluorescence often have tunable fluorescence when coordinating with central metal atoms.\(^\text{162,163}\) ICP-type nanomedicines are believed to be suitable candidates for computational design and combinatorial chemistry for drug content-specific nanomedicine development, because of their high drug-loading content and efficiency.\(^\text{164}\) This section outlines recent advances in ICP I-type high drug-loading nanomedicines that consist of metal atoms and two kinds of organic ligands, with one of the ligands being the drug molecule.

The first use of ICPs in drug-delivery systems was in 2011 by Xing et al.\(^\text{164}\) They defined ICP I-type nanomedicines as “host metal–ligand coordination polymer” and ICP II-type nanomedicines as “metal–ligand coordination polymer”. As illustrated in Figure 7, in ICP I-type nanomedicine, host ligands and drug molecules were coordinated with metal ions. For some “metal–drug” coordination particles with unsuitable bond energy or irregular NP morphology, host ligands, such as PEG, oligochitosan, and Pluronic F127, were introduced to endow the particles with appropriate size and pH sensitivity. For example, mitoxantrone (MX)-Cu NPs did not release MX, even in a highly acidic medium, because of their strong coordination bond. In this case, PEG was introduced as the host molecule to weaken the coordination bond, which was achieved based on the weak coordination binding site provided by the EO moiety of PEG. Although the article did not provide a specific drug-loading content, such values were obviously high in these nanomedicines. With this strategy, Moustaoui et al reported another ICP I-type high drug-loading nanomedicine, PEG-Au(III)-Dox, in a recent work.\(^\text{165}\) In this nanomedicine, Dox molecules and dicarboxylic PEG molecules were chelated with Au(III) ions from tetrachloroauric acid (HAuCl\(_4\)). It was demonstrated that nanomedicine depolymerization and effective Dox release were achieved at pH 4, with stability attained at a diameter close to 20 nm in physiological pH. The drug-loading efficiency of Dox in this nanomedicine was up to 85%.

The following issues are worth noting. This strategy is only applied to drugs possessing complexation ability, such as carboxylates and phosphonates, including daunorubicin

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**Figure 7** Formation of ICP I-type nanomedicine and ICP II-type nanomedicine and their pH-responsive release. **Abbreviation:** ICP, infinite coordination polymer.
hydrochloride, Dox hydrochloride, MX, alizarin red, 1,10-phenanthroline, cis-platinum, 1,4-bis-([2-{dimethylamino-}
N-oxide}ethyl]amino)5,8-dihydroxyanthracene-9,10-dione (AQ4N), and gossypol. The particle size and pH-responsive point of nanomedicines are dependent on the complex combinations of host ligands, drug molecules, and center metal ions. Computational design and combinatorial chemistry can be utilized as primary screening to equip ICP I-type nanomedicines with ideal properties. More studies on cell and animal levels are needed to confirm the outstanding antitumor effects of ICP I-type nanomedicines.

**Carrier-free nanomedicines**

With the previously mentioned strategies, a significant number of high drug-loading nanomedicines have been developed (Tables 1 and 2). However, an intrinsic problem in these nanomedicines is that their degradation and excretion rates remain undetermined. To circumvent this concern, carrier-free nanomedicines with no excipient were developed, such as traditional DNCs, DDC micelles, MBioFs, and ICP II-type nanomedicines. All these types of nanomedicines often have drug-loading contents higher than 80 wt%. These strategies are summarized in Table 3. Obviously, these strategies were applied for nanomedicine loading with drugs that are stable in the circulatory system, which are similar to those shown in Table 2.

**Drug nanocrystals**

DNCs are a conventional viable strategy to develop hydrophobic drug nanomedicines with the advantages of general applicability and simplicity. Basically, such nanomedicines consist of DNCs dispersed in liquid medium, and are stabilized by surface-active agents, such as the poloxamer family, Tween, and polymers. For oral administration, the high surface:volume ratios of DNCs significantly increase dissolution velocity, thereby leading to an improvement in the gastrointestinal tract absorption based on the Noyes–Whitney equation. For intravenous administration, the smallness of this nanomedicine enables it to enter blood circulation and achieve targeted delivery through further surface modification. In general, the conversion of a medical technology converted into a market product takes several decades, but a DNC-type nanomedicine (also called nanosuspensions) only needs several years. The first DNC product for oral application (named Rapamune®) was successfully introduced to the market by Wyeth in 2000. Moreover, the first DNC product for intravenous application (named Abraxane®) was placed on the market by American Pharmaceutical Partners. Currently, several products are undergoing clinical trials and will be on the market in the near future. DNC-fabrication techniques are classified as either “top-down” or “bottom-up” methods, based on the size-change process. The top-down method starts from large crystals to small NCs in a step-by-step manner during a high-energy process, such as pearl milling and high-pressure homogenization. The bottom-up method involves procedures that start from molecules to NCs, such as solvent exchange, high-gravity-control precipitation, evaporative precipitation, rapid expansion of supercritical solution, and supercritical antisolvent methods. DNCs prepared via bottom-up methods need less energy and are smaller compared with those prepared through top-down methods. Solvent exchange is the simplest, commonest, and most cost-effective method used in laboratory experiments. This method can produce both crystalline and amorphous forms of DNCs. DNCs smaller than 200 nm can be administered intravenously with 100% bioavailability and achieve multifunctionality. Li et al developed differently shaped 10-hydroxy-CPT (HCPT) NCs, nanospheres (NSs), and nanorods using solvent exchange and further surface modification. Polymaleic anhydride-alt-1-octadecene–PEG (C18PMH-PEG) was chosen as a stabilizer, which was

**Table 3** Overview of three strategies of carrier-free nanomedicines

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Structure diagram</th>
<th>Main drug-loading mechanism</th>
<th>Application range</th>
<th>Drug-loading content</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drug nanocrystals</td>
<td><img src="image" alt="Structure diagram" /></td>
<td>Crystallization</td>
<td>Hydrophobic drugs mostly</td>
<td>≤93.3 wt%</td>
<td>21, 169–180, 182–200</td>
</tr>
<tr>
<td>Amphiphilic drug-drug conjugate</td>
<td><img src="image" alt="Structure diagram" /></td>
<td>Covalent bond</td>
<td>Hydrophobic–hydrophilic drug pairs</td>
<td>≤100 wt%</td>
<td>201–204</td>
</tr>
<tr>
<td>MBioFs</td>
<td><img src="image" alt="Structure diagram" /></td>
<td>Coordinate bond</td>
<td>Drugs with complexing ability (both hydrophobic drugs and hydrophilic drugs)</td>
<td>≤75 wt%</td>
<td>207</td>
</tr>
<tr>
<td>ICP II-type</td>
<td><img src="image" alt="Structure diagram" /></td>
<td></td>
<td></td>
<td>≤79.1 wt%</td>
<td>209–213</td>
</tr>
</tbody>
</table>

**Abbreviations**: CP, coordination polymer; MBioFs, metal–biomolecule frameworks; ICP, infinite coordination polymer; wt, weight.
introduced on the surfaces of DNCs through noncovalent hydrophobic interaction. The surface PEGylation on HCPT NCs served as a steric repulsion–hydration molecular layer to reduce electrostatic and hydrophobic interactions with biomolecules, endowing them with excellent long-term stability at high salt concentrations and extreme pH. The drug-loading contents of PEGylated NSs and nanorods were as high as 91.9 wt% and 93.3 wt%, respectively. In other research, HCPT NCs with targeted delivery ability were prepared using solvent exchange, followed by surface functionalization with C18PMH-PEG-Fa. This nanomedicine benefited from the high selectivity of Fa to folate receptor-positive cancer cells, resulting in enhanced drug efficacy. The drug-loading content in this nanomedicine was approximately 78%. DNCs can also be used for the fabrication of an all-in-one processing system for cancer diagnosis and treatment. Zhou et al developed carrier-free multifunctional multi-DNCs (MDNCs) that consisted of methotrexate (Mtx), HCPT, Ptx, and the amphiphilic polymer stabilizer C18PMH-PEG (Figure 8). The MDNCs were prepared through solvent exchange, and the PEG layer was added on the surface via hydrophobic interaction. In vitro studies demonstrated that the MDNCs showed synergistic effects and improved tolerance. Given the minute amount of red fluorescent dye (TBPT, synthesized by themselves), the MDNCs exhibited high contrast during in vivo imaging. The drug-loading contents of Mtx, HCPT, and Ptx were 24.3 wt%, 49.1 wt%, and 26.6 wt%, respectively.

DNCs are a traditional assembly technology utilized in the fabrication of high drug-loading nanomedicine. Considerable developments have been made in NC-assemble technology over the last decade. However, certain limitations still exist: 1) both top-down and bottom-up methods apply only to the fabrication of hydrophobic high drug-loading nanomedicines, and thus the range of application is limited by the requirement of drug solubility; 2) the problem of removing residual organic solvent cannot be neglected for industrial production; and 3) most studies on crystal nanomedicine have focused on the solubility problem of hydrophobic drugs, and few have involved sustained release, targeting problems, and cell uptake of DNCs. Therefore, research on surface modification and cyclic behavior of nanomedicine crystal in vivo represents an important direction for further DNC development.

Amphiphilic drug–drug conjugates

ADDCs are a class of special PDCs in which the hydrophilic and hydrophobic parts are represented by hydrophilic and hydrophobic drugs, respectively. This class of dual-drug-loaded nanomedicines has the advantages of ultrahigh drug-loading content and synergistic combination chemotherapy because of its inherent nature. This strategy, which was first proposed by Huang et al., is considered an excellent candidate for the fabrication of dual-drug synergistic therapy nanomedicines with high and fixed drug loading. Two methods are proposed under this strategy. The first is the “natural method,” which is applied to drug pairs when one drug has a carboxyl group and the other an activated hydroxyl group. The second is the “modified method,” which is applied to any modifiable AD pairs using a linker compound. All these nanomedicines have sizes suitable for intravenous administration. The total drug-loading content is approximately 100 wt%.

For the natural method, a dual-drug-loaded nanomedicine was synthesized from a hydrophilic anticancer drug, irinotecan (Ir), and a hydrophobic anticancer drug, chlorambucil (Cb), via a hydrolyzable ester linkage by Huang et al. The ester linkage was formed from the hydroxyl of Ir molecules and carboxyl of Cb molecules through direct esterification using dicyclohexylcarbodiimide/4-dimethylamino-pyridine without any addition (Figure 9A). The amphiphilic Ir-Cb conjugates self-assembled into micelles in aqueous solution with average size of approximately 75.7 nm. After cell uptake of nanomedicine, the ester linkages were hydrolyzed by the existing esterase and low pH in endosomes, causing the drug molecules to be effectively released. In vitro and in vivo antitumor experiments demonstrated that this nanomedicine exhibited high anticancer activity and could circumvent the multidrug resistance of tumor cells in chemotherapy. The same method was also used to fabricate fluoroxyuridine–bendamustine (FUDR-Bdm) and Ir-Bdm conjugates by Huang et al.

The range of application of this strategy is widely extended with the use of the modified method. For example,
in a recent work of Hu et al., a dual-drug-loaded nanomedicine, CPT-FUDR conjugates, was synthesized through two-step esterification of hydrophobic CPT and hydrophilic FUDR using a linker compound. As illustrated in Figure 9B, the first step was the introduction of a carboxylic group onto the CPT molecule through esterification of CPT and diglycolic anhydride. Subsequently, the esterification reaction of CPT-COOH and FUDR was carried out, and the amphiphilic CPT-FUDR conjugate linked by a hydrolyzable ester was formed. The resultant amphiphilic conjugate self-assembled into nanomicelles with hydrodynamic size of 36.4 nm. In vitro studies demonstrated that CPT-FUDR conjugates exhibited synergistic anticancer efficacy and similar intracellular behavior to DDC nanomedicines prepared via the natural method.

The following findings are noted. ADDCs are a kind of high drug-loading nanomedicine with narrow applicable range. This requires one of the drugs to be hydrophilic and the other to be hydrophobic, and these two drugs should satisfy a specific structure simultaneously. Few drug pairs possess these conditions while having a synergistic effect at the same time. Although some ADDC pairs have been reported to possess good anticancer efficacy, the influence of the structural modification of drug molecules on anticancer efficacy and pharmacokinetics is impossible to predict, especially the influence of the structural transformation of drug molecules caused by linker compound in the modified method. Further surface modification is difficult, due to the micelle-type structure of ADDC nanomedicines. Therefore, additional research should be concentrated on the surface modification of drug molecules and pharmacokinetics in the future.

Coordination polymer NPs

In this section, we focus on MBioFs and ICP II-type high drug-loading nanomedicines, in which the drug molecule acts as the only ligand coordinating with the central metal atom without any other vehicle components. Similar to ICP I-type nanomedicines, MBioFs and ICP II-type strategies are only applicable to those drugs with complexing ability.

MBioFs

MBioFs are defined as MOFs constructed from biomolecules that serve as organic ligands, such as amino acids, peptides, proteins, nucleobases, saccharides, and drug molecules (Figure 4). MBioFs display more advantages in the fabrication of high drug-loading nanomedicines than MOFs: 1) MBioFs avoid the structural requirements of MOFs, such as pore size and volume; 2) drug release is achieved through nanostructure biodegradation, without any side effects,
because no inert material linkers are used; and 3) drug molecules often have many different metal-binding sites, which leads to multiple possible coordination modes and adjustable physical and chemical properties. For example, Miller et al developed a small-pore iron (II/III) nicotinate Bio-MIL-1, which consists of nicotinic acid (pyridine-3-carboxylic acid, also called niacin or vitamin B₃) and nontoxic iron. The structure of Bio-MIL-1 comprises a three-dimensional connected framework built from trimeric Fe₃N₅O₁₃ units linked together via nicotinate molecules. These nanomedicines have a drug-loading content as high as 75 wt%, and are rapidly degraded to achieve nicotinic acid release under physiological conditions. To date, MBioF nanomedicines have not been fully explored. Only some examples have been reported, and almost no studies on cell and animal levels can be searched. A possible explanation is that the stability of some MBioFs in aqueous and in vivo environments is not yet known.

As such, further in vitro and in vivo studies should be conducted so that MBioFs have wide application prospects in the biomedical field.

**ICP II-type nanomedicines**

ICP II-type nanomedicines are ICPs prepared directly from drug molecules and metal atoms (Figure 7). ICP II-type nanomedicines display the same advantages as ICP I-type nanomedicines, and have higher drug-loading content. ICP II-type nanomedicines can be formed from a number of self-assembly processes. Solvothermal synthesis is the most straightforward method, and consists of metal and ligand molecules heated in a solvent, which can lead to the formation of amorphous or crystalline products. ICP II-type can also be synthesized using reverse microemulsion (water in oil), in which the reactants are suspended in surfactant-stabilized aqueous droplets in the organic phase and the droplets behave as nanoreactors, facilitating ICP assembly. Finally, ICPs can be formed via rapid precipitation that involves precipitating a solution of reactants with a solvent, which usually results in amorphous particle formation. For instance, Rieter et al fabricated an ICP II-type nanomedicine from Tb(III) ions and \( \text{c,c,t-} (\text{PtCl}_2[\text{NH}_3]_2[\text{O}_2\text{C}\text{CH}_2\text{CH}_2\text{CO}_2\text{H}]) \) (disuccinatocisplatin, [DSCP]) via rapid precipitation (Figure 10).

The Tb(III)-DSCP ICPs displayed a size of 58.3±11.3 nm in diameter using dynamic light scattering. The drug-loading content of DSCP was 36 wt%, which was determined through thermogravimetric analysis. These ICP particles were easily functionalized with a variety of silyl-derived molecules and further grafted to silyl-derived c(RGDfK), which is a small cyclic peptide sequence that is frequently observed in many angiogenic cancers, on the surface to achieve targeted delivery. Huxford et al synthesized serious ICP II-type Mtx nanomedicines via reverse microemulsion. Different metal ions, such as Zn²⁺, Zr⁴⁺, and Gd⁴⁺, with different coordination numbers were chosen to be coordinated with Mtx. The ICPs were further stabilized and modified using dioleoyl trimethylammonium propane/dioleoyl-\( \text{L-} \alpha\)-phosphatidylethanolamine–anisamide. These nanomedicines were spherical in appearance with diameters between 40 and 100 nm via transmission electron microscopy. Thermogravimetric analysis revealed that the drug-loading content of Mtx reached 79.1 wt%.

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**Figure 10** Synthesis and functionalization of Tb(III)-DSCP ICPs.  

**Abbreviations:** DSCP, disuccinatocisplatin; ICPs, infinite coordination polymers; NCP, nano-coordination polymer; PVP, polyvinylpyrrolidone; TEOS, tetraethyl orthosilicate.
MBioF and ICP II-type strategies are not broad-spectrum techniques, due to the requirement of drug-complexing ability, similar to ICP I-type strategy. In addition, controlling the size of these nanomedicines is not easy, because of the absence of host ligands. Fortunately, molecular simulations are possible, because of the one-component feature and ultrahigh drug-loading content and efficiency of these nanomedicines. The application of computational design and combinatorial chemistry will allow the efficient design of MBioF and ICP II-type high drug-loading nanomedicines.

**Nanomedicines following niche and complex strategies**

Some niche and complex strategies, such as aqueous noncovalent assembly and multiple assemblies, are used for the fabrication of specific multidrug-loaded high drug-loading nanomedicines (Table 4). For aqueous noncovalent assembly, the selected drugs should possess opposite static electricity, $\pi-\pi$ stacking, and hydrophobic interactions. The biggest difference between solvent exchange for DNC fabrication and noncovalent assembly is that the former is applied for the fabrication of hydrophobic drug-loaded nanomedicines, whereas the latter is utilized for the fabrication of hydrophilic (including water-soluble drugs under certain conditions) drug-loaded nanomedicines. In this strategy, no organic solvent is used in the fabrication process. The molar ratio of multiple drugs is tailorable within a certain range, and the total drug-loading content for nanomedicines is 100 wt%.

Zhang et al developed a carrier-free, chemophotodynamic dual-drug-loaded nanomedicine prepared from a chemotherapeutic agent (Dox) and a photosensitizer (chlorine $\text{e}_6\ [\text{Ce}_6])$. Depending on electrostatic, $\pi-\pi$ stacking, and hydrophobic interactions, drug molecules self-assembled into well-defined NSs with an average size of 70 nm in a basic aqueous solution (pH 12). Total drug-loading content was 100 wt%, and drug-loading efficiency of $\text{Ce}_6$ and Dox were 95% and 99%, respectively. Further experiments on BALB/c nude mice with xenografted MCF7 tumors demonstrated that this carrier-free, chemophotodynamic dual-drug-loaded nanomedicine was an effective combinational therapeutic modality. Similarly, a novel high drug-loaded nanomedicine is currently being studied by our group.

For multiple assemblies, the selected drug–carrier pairs should possess appropriately matched physical and chemical properties. Notably, this strategy is a comprehensive strategy for the fabrication of multidrug-loaded nanomedicine. Recently, our group reported a mulberry-like dual-drug-loaded nanomedicine following this strategy. First, two perfectly matched drug–carrier pairs, apogossypolone–cationic amphiphilic starch ($\text{ApoG}_2-\text{CSaSt}$) and Dox-HA, self-assembled to form two seeds, namely, $\text{ApoG}_2-\text{CSaSt}$ micelles and Dox-HA NSs (DHA NSs), respectively. These two seeds further self-assembled to form NPs (MLDC NCs) with a mulberry-like shape and dynamic size of 83.1±6.6 nm. The first self-assembly occurred based on the hydrophobic interaction in the $\text{ApoG}_2-\text{CSaSt}$ pair and the electrostatic absorption in the DHA pair. The subsequent self-assembly occurred based on the electrostatic absorption between CSaSt and HA caused by the designed positive charge of CSaSt and the natural negative charge of HA. DHA NSs located on the surface of MLDC NCs functioned as a targeting agent at the same time. The drug-loading contents of $\text{ApoG}_2$ and Dox in MLDC NCs were 13.3±1.2% and 13.1±3.7%, respectively, and these values were adjustable to a certain extent. For the in vivo tumor-suppression test, the injection dosages of Dox and $\text{ApoG}_2$ were both 2 mg/kg, which was only a fifth of the normal dosages in the mouse test, and resulted in enhanced antitumor effects. This result suggested that high drug-loading nanomedicines can effectively realize multidrug synergy and have obvious antitumor advantages compared with low drug-loading nanomedicines.

Deng et al developed an anticancer drug and siRNA-loaded nanomedicine following a similar strategy. First, a Dox-loaded negatively charged phospholipid liposome was developed. Consequently, a siRNA-loaded film was generated through alternate deposition of positively charged poly-L-arginine and negatively charged siRNA atop the Dox-loaded liposome. Finally, the NP surface was coated with a layer of HA to achieve targeted delivery. The loading efficiency and content of Dox in the nanomedicine were

<table>
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<th>Table 4 Overview of two strategies used for minority-specific nanomedicines</th>
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<td>Strategy</td>
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<td>Noncovalent assembly</td>
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<td>Multiple assembly</td>
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Abbreviation: wt, weight.
97 wt% and 5.5 wt%, respectively. In addition, the poly-L-arginine/siRNA layer-by-layer film loaded approximately 3,500 siRNA molecules per NP per layer. Niche and complex strategies are usually applicable for particular drugs or drug pairs, and their implementation schemes are unsuitable for other nanomedicines. However, the design philosophy, which ingeniously takes advantage of the special physicochemical property of drugs or drug pairs, is worth learning from.

Conclusion and prospects
In the past 30 years, the development of nanocarrier-based platform technologies has led to significant progress in nanomedicine fabrication and applications. However, most of the existing nanomedicines have low drug-loading content (generally lower than 10%), causing extra system toxicity and burden on patients to excrete carrier materials. Therefore, it is difficult for most of the carrier-based nanomedicines to get FDA approval, which limits their applications in the clinic. With this in mind, emergence and fabrication of high drug-loading nanomedicines may not provide the final answer, but present a promising alternative approach to solve this problem, because of the reduced or avoided use of carrier materials.

The three classes of high drug-loading nanomedicines, which can either reduce or avoid the use of carrier materials, are applied to the fabrication of high drug-loading nanomedicines with different drug properties. In addition to common advantages of nanomedicines, some high drug-loading-nanomedicines possess inherent advantages, benefiting from their structure and ultrahigh drug-loading content and efficiency, such as stimulant responsibility, multidrug combined therapy, computational design, and combinatorial chemistry properties. Despite all these benefits, the therapeutic effect of current high drug-loading nanomedicines is not so satisfactory. Furthermore, several theoretical and actual problems remain unsolved. Demonstrating the better in vivo antitumor effect of high drug-loading nanomedicines than low drug-loading nanomedicines is significant. However, most existing high drug-loading nanomedicine studies have focused on design and fabrication and lacked comparison of the antitumor effect at the animal level. As such, these studies failed to demonstrate the complete advantages of high drug-loading nanomedicines. Understanding the differences between the antitumor mechanisms of high drug-loading and low drug-loading nanomedicines is important to improve the design of subsequent high drug-loading nanomedicines. However, existing reports on high drug-loading nanomedicines have mostly focused on nanomedicine design, and almost no research has been conducted on the internalization mechanism, intracellular release, and subcellular level actions. Owing to long conversion times, safety concerns, and associated socioeconomic uncertainties, most concepts investigated are at the laboratory stage, and a few have entered routine clinical application, such as Rapamune and Abraxane, which we discussed in this paper. Despite this, it irresistible that more high drug-loading nanomedicines will be transitioned into clinics in the future.216

Meanwhile, the following issues are worth noting. For some specific drugs, such as gene drugs that are unstable in the circulatory system and brain-targeting drugs that need to cross the blood–brain barrier, carrier materials are essential for the realization of their function. For some carrier materials with function, such as Pluronic block copolymers, biological response modifiers affecting cell-membrane microviscosity and ATPase activity of drug-efflux transporters, the use of carrier materials will enhance the therapeutic effect.217 For most of the high drug-loading nanomedicines, it is hard to realize in vivo controlled release, such as ultrasound-controlled release, magnetism-controlled release, and microwave-controlled release. For nanomedicines with drug-loading content higher than 50%, sustained-release effects are far below expected, and there are fewer pharmacokinetic studies in this category of articles. For nanomedicines with drug-loading content higher than 90%, it is usually hard to achieve surface modification, so further study is needed on the action of those high drug-loading nanomedicines in the complicated physiological environment and highly dynamic and heterogeneous tumor sites. Clearly, the high drug-loading property serves therapeutic effect and function purposes.

Over the past few years, the number of FDA-approved drugs, including protein drugs, oligonucleotide drugs, and small-molecule inhibitors, has increased, such as lixisenatide,218 defibrotide sodium,219 and rucaparib.220 Although these drugs show excellent performance against certain diseases, it cannot be ignored that their inherent drawbacks, such as poor water solubility and unstable properties in the circulatory system, lead to unsatisfactory absorption and frequent injection for patients. A promising approach to overcome these drawbacks is loading them into nanomedicine. Even though much time and high costs are required to develop a new nanomedicine, we still believe that there will be more nanomedicines approved by the FDA in the future. We hope that this review will attract more attention from researchers in this field, and expect that high drug-loading nanomedicines with still-better properties will be prepared by developing novel carrier materials, exploring new fabrication strategies and modification methods and perfecting theoretical research at the cell and animal level. We also believe
that high drug-loading nanomedicine-delivery systems will be gradually improved through the joint efforts of chemists, biologists, material specialists, and cancer and pharmaceutical researchers, and will occupy an important position in the field of drug-delivery systems in the future.

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**Disclosure**

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