Tumor-targeting delivery of herb-based drugs with cell-penetrating/tumor-targeting peptide-modified nanocarriers

Abstract: Cancer has become one of the leading causes of mortality globally. The major challenges of conventional cancer therapy are the failure of most chemotherapeutic agents to accumulate selectively in tumor cells and their severe systemic side effects. In the past three decades, a number of drug delivery approaches have been discovered to overwhelm the obstacles. Among these, nanocarriers have gained much attention for their excellent and efficient drug delivery systems to improve specific tissue/organ/cell targeting. In order to enhance targeting efficiency further and reduce limitations of nanocarriers, nanoparticle surfaces are functionalized with different ligands. Several kinds of ligand-modified nanomedicines have been reported. Cell-penetrating peptides (CPPs) are promising ligands, attracting the attention of researchers due to their efficiency to transport bioactive molecules intracellularly. However, their lack of specificity and in vivo degradation led to the development of newer types of CPP. Currently, activable CPP and tumor-targeting peptide (TTP)-modified nanocarriers have shown dramatically superior cellular specific uptake, cytotoxicity, and tumor growth inhibition. In this review, we discuss recent advances in tumor-targeting strategies using CPPs and their limitations in tumor delivery systems. Special emphasis is given to activable CPPs and TTPs. Finally, we address the application of CPPs and/or TTPs in the delivery of plant-derived chemotherapeutic agents.

Keywords: cancer, nanocarriers, cell-penetrating peptide, targeting drug delivery, herb-based drug, tumor targeting

Introduction

Despite great advancements in novel drug delivery systems, cancer nevertheless remains one of the main causes of morbidity and mortality worldwide. In 2015, cancer was accountable for 8.8 million deaths. Globally, for every six deaths, one is due to cancer. Within the coming two decades, this is estimated to increase by about 70%. In addition, the economic burden due to cancer has been significantly increasing. In 2010, the overall annual economic cost of cancer was estimated at approximately US$1.16 trillion.¹

Chemotherapy, radiation, and surgery with chemotherapy are the major treatment protocols for cancer. Utilization of chemotherapy has been demonstrated to enhance the survival rate of patients with malignant cancer slightly. Surgery and radiotherapy are the best treatments for local tumors and nonmetastatic cancers, and they are also helpful in cancer that has not been disseminated throughout the body. Therefore, chemotherapy is the treatment of choice for metastatic malignancies, since they are well distributed to every organ in the body.² However, most conventional chemotherapeutic agents...
currently in clinical use are limited, due to several undesirable properties, including poor solubility and bioavailability, rapid elimination from systemic circulation, narrow therapeutic index, and unselective site of action after intravenous/oral administration and cytotoxicity to normal tissues, which can be the main obstacles to success of cancer therapy.\(^3\)

Furthermore, chemotherapeutic agents are often unable to penetrate and reach the internal parts of solid tumors, leading to inefficient cytotoxicity to cancerous cells.\(^4\) The other limitation of traditional chemotherapy is associated with Pgp, a multidrug-resistant (MDR) protein that is overexpressed in cancer cells, acts as an efflux pump and decreases accumulation of drug inside the tumor, and plays a key role in the development of resistance to anticancer drugs. Therefore, the drugs administered remain unsuccessful in exerting the desired effect.\(^3,5\) Though different drug delivery approaches consisting of antibodies, hormones, and growth factors have been designed and administered, significant internalization the cancerous cells cannot be achieved, due to the reticuloendothelial system and intracellular enzymes.\(^6\) Consequently, in recent decades, much work has been devoted to overcoming the major drawbacks of conventional cancer chemotherapy.

Nanocarrier-based drug delivery systems have been shown to have great potential in reducing the aforementioned problems of cancer therapy.\(^7\)–\(^13\) Moreover, decorating nanocarrier surfaces with different ligands targeting specific receptors that are overexpressed on tumor cells has been exploited for active targeting of chemotherapeutic agents to cancer cells.\(^14\)

Most importantly, nanocarriers functionalized with cell-penetrating peptides (CPPs) and/or tumor-targeting peptides (TTPs) have been a highly promising strategy and attracted the attention of researchers. These peptides are very advantageous, as they efficiently deliver a broad variety of cargo intracellularly and extracellularly.\(^15\)–\(^17\) CPP-mediated drug delivery is achieved either by the formation of stable, noncovalent complexes or by a covalent bond with the cargo.\(^16\) However, most of the first-generation CPPs suffered from a lack of cell-type specificity and in vivo stability, which led to the emergence of activable CPPs (ACPPs) and TTPs. In this review, we focus on recent advances in CPPs and TTPs, and the tumor-targeting approach of herb-based anticancer bioactive substances using CPP- and/or TTP-modified nanocarriers.

**Overview of nanocarrier-based tumor-targeting drug delivery approaches**

Targeted cancer therapy is viewed as an irreplaceable component of current anticancer drug development.\(^18\) The best strategy to improve the efficacy and reduce the toxic effects of an anticancer drug is delivering it to a specific target and keeping sufficient concentrations for enough time, which allows the drug to produce the desired therapeutic effect. Cancer cells can be targeted by two approaches: passive targeting and active targeting.

Passive targeting involves the extravagation of drug formulations through leaky vasculature/capillaries of tumors that result from abnormal angiogenesis at sites of tumors, resulting in accumulation and retention. This phenomenon is known as the enhanced permeability and retention effect\(^3,19\) (Figure 1). In addition, the release of drug from pH-sensitive formulations in the acidic microenvironment inside the...
cancer cell,\textsuperscript{6,20} phagocytosis of the particulate carrier by mononuclear phagocytosis systems, and privileged localization in the organs of the reticuloendothelial system are also considered passive targeting.\textsuperscript{21}

Nanoparticle (NP)-based drug delivery plays a crucial role in the passive targeting of tumors.\textsuperscript{22} NPs possess unique properties, such as large surface area, small size, high efficiency in encapsulating a wide variety of drugs, and modifiable surface chemistry. These benefits make NPs a potentially superior treatment approach to traditional cancer therapies. The size of drug carriers and the abnormal and permeable vasculature of tumors are the base for passive targeting.\textsuperscript{23} Most tumors manifest an abnormally dense and leaky vasculature formed via stimulation by VEGF. In normal vasculature, particles >2 nm are prevented from crossing through endothelial cells due to tight junctions.\textsuperscript{24} However, in tumor vasculature, tight junctions and the basement membrane are disordered, allowing passage of particles of 10–200 nm through the leaky neovasculature of the tumor and then retention in the tumor. Furthermore, the poor venous and lymphatic system in the tumor creates an opportunity for the NPs to accumulate therein at high concentrations for a long time.\textsuperscript{3} The size and surface charge of NPs are known to affect the half-life and biodistribution of NPs significantly. Larger NPs (>100 nm) are usually cleared from the circulation by phagocytosis. Similarly, very small NPs (<10 nm) have a high rate of clearance. The positive charge on the surface of particles is known to assist internalization into cancer cells. Moreover, surface modification of NPs with such polymers as polyethylene glycol (PEG) can increase the circulation time of particles and increase the accumulation of NPs at tumor sites but can affect cellular uptake by cancer cells.\textsuperscript{25}

In addition to targeting potential, nanocarriers offer protection of the drug from degradation, decrease renal elimination, increase its half-life in the circulation, allow controlled release kinetics, and improve solubility. Furthermore, NPs have known to have the potential to bypass MDR mechanisms by various approaches.\textsuperscript{2,10,26}

Several types of nanocarrier have emerged for the drug delivery of chemotherapeutic agents, including magnetic and metallic NPs, such as iron oxide or gold NPs, silver NPs, nanodiamonds, carbon-based structures (graphene sheets and carbon nanotubes), polymeric NPs, dendraimers, quantum dots, hydrogel-based delivery systems, silica-based NPs, lipid-based NPs (liposomes), solid-lipid NPs, nanostructured lipid carriers (NLCs), viral NPs, and hybrid NPs.\textsuperscript{27–31}

Passive targeting of almost all nanocarriers occurs with poor selectivity and insufficient tumor cell uptake. A more advanced approach of targeting for oncology applications is the modification of the surface of NPs with specific tumor-homing ligands.\textsuperscript{32} The ligands are known to bind to receptors that can be overexpressed on the surface of a cancer cell. Ligands with selective affinity for a specific receptor or molecule differentially expressed at the target site are presented on the surface of nanocarriers, resulting in selective accumulation and cellular uptake at the site of action.\textsuperscript{33} This strategy significantly increases accumulation and retention of NPs in the tumor vasculature and specific and successful internalization by target tumor cells, which is known as active tumor targeting.\textsuperscript{34} Ligands that have been used to modify NPs include monoclonal antibodies, folate acid, hyaluronic acid, albumin, vitamins (folate, vitamin B\textsubscript{12}, thiamine, and biotin), transferrin, lectins, aptamers, and peptides.\textsuperscript{6,21,23,35}

Most recently, peptide-based targeting ligands have been attracting the attention of researchers and are the main concern of this review. CPPs provide several advantages over other ligands.\textsuperscript{36} Peptide ligands are highly selective for target tissue/cells, and multiple ligands can be conjugated to a single drug carrier to offer multivalent conjugation, thus increasing binding affinity to the target.\textsuperscript{37} CPP details are presented in the following section.

### CPP/TTP-based drug delivery

#### CPPs and classification

A recent advancement in the field of molecular biology that appears to have had an enormous impact on cancer therapy is the innovation of the tissue- or cell-penetration system.\textsuperscript{15,38} CPPs are short peptides that are able to cross through tissue and cell membranes in an energy-dependent or -independent manner and are used to transport a wide variety of bioactive conjugates (cargoes), including proteins, peptides, DNA, siRNA, small drugs, fluorescent compounds, NPs, and other substances, into cells.\textsuperscript{39} CPPs are extremely beneficial, because they are biocompatible and the peptide sequence can be altered to fine-tune hydrophobicity, affinity, charge, solubility, and stability. They can also be readily synthesized in sufficient quantity.\textsuperscript{17}

Besides the use of CPPs as inert vectors for the transport of cargo molecules, dual-acting CPPs that have properties of both cell permeation and bioactivity have been emerging. Selected CPPs are known to produce bioactivity, including neuroprotecting effects,\textsuperscript{40} induce apoptosis in cancer cells,\textsuperscript{41,42} and suppress breast tumorigenesis.\textsuperscript{43} More recently, a CPP obtained from azurin (p28) was found to prevent
phosphorylation of VEGFR2, FAK, and Akt, leading to inhibition of tumor growth and angiogenesis.\textsuperscript{44}

CPPs can be conjugated to cargoes by either noncovalent complex formation or covalent bonds. Covalent conjugation of a CPP may be obtained chemically through disulfide bonds, amine bonds, or specific linkers that enable the release of the cargo when internalized into the cell.\textsuperscript{42} However, the possibility of changing the bioactivities of the conjugates is the main risk of the covalent conjugation of CPPs.\textsuperscript{45} Furthermore, covalent methods are associated with such problems as lack of suitable reactive groups on the polymer, unstable intermediates, and inefficient coupling and purification. In these cases, the noncovalent approach seems more appropriate.\textsuperscript{46} The noncovalent complex is formed by electrostatic and/or hydrophobic interactions between positively charged CPPs and negatively charged drug carriers.\textsuperscript{47}

CPPs can be classified by their origin, function, sequence, or mechanism of uptake. According to their physicochemical properties, CPPs can be cationic, hydrophobic, or amphiphilic. Cationic CPPs have highly positive net charges at physiological pH. They originate from the basic short strands of arginines and lysines, eg, TAT\textsubscript{48-60} (GRKKRRQRRRPPQ), penetratin (RQIKIKWFQNRRMKWKK), and DPV1047 (VKRGLKLRHVRPRVTRMDV). Hydrophobic CPPs primarily contain apolar residues, which have amino-acid groups that are vital for cellular uptake, along with a low net charge, eg, Pep7 (SDLWEMMMVSLACQY) and C105Y (CSIPPEVFKNKFVYLI) are some of the hydrophobic peptides. The third class of CPPs is the amphiphilic CPPs, which contain both hydrophilic and apolar hydrophobic regions of amino acid sequences. Pep1 (KETWWETWTEWSQPKKRKV) and pVEC (LLIIILRRRIRKQAHAHSK) are typical examples. Based on their origin, peptides can be classified as derived CPPs, chimeric CPPs, and synthetic CPPs.\textsuperscript{48-49} Detailed reviews on CPP classification can be found elsewhere.\textsuperscript{39,49,50}

**Limitations of CPPs**

Despite the great potential of CPPs as carriers for drug delivery to cancer, most conventional CPPs suffer from a number of limitations, including lack of cell specificity and in vivo instability, which are the main obstacles to their further clinical development.\textsuperscript{51}

**Lack of cell specificity**

In vivo applications of CPP are limited because of their aspecific tissue/cell penetration. To date, several approaches have been suggested to target cancer cells selectively using CPPs. Among these, ACPPs and the combined use of CPPs with TTPs are discussed in the following sections.

**In vivo degradation**

In vivo stability should be taken into account for CPPs to be beneficial as vectors for the delivery of any therapeutic agent. Due to their peptidic nature, CPPs are degraded both extracellularly and intracellularly by proteases.\textsuperscript{52,53} In vivo CPP stability depends on factors that influence proteolytic cleavage, including amino acid sequence, conformation, chemistry (unnatural amino acids, D-isomers, chemical bridges, amino acid modifications), routes of administration, type of conjugate cargo, and method of conjugation of CPPs with this cargo.\textsuperscript{53}

Several strategies have been employed to improve the in vivo stability of CPPs. One approach is the synthesis of retro-inverse (RI) CPPs, introduced by Chorev and Goodman.\textsuperscript{54} RI CPPs are peptides consisting in D-amino acids in the reverse sequence of the naturally occurring L isoforms. Subsequently, RI transformation has extensively been used as an approach of producing a proteolytic resistant peptide analogues.\textsuperscript{55} However, RI peptides are reported to cause cytotoxicity, decline in metabolic activity, morphological changes, and induction of apoptosis.\textsuperscript{56} In addition, end-to-end cyclization of CPPs have been proposed to attain improved metabolic stability. End-to-end cyclization is known to offer greater proteolytic resistance and higher internalization efficiency as a result of improved structural or conformational rigidity/stability. Cyclization through an intramolecular disulfide bond has also been reported to improve in vivo stability.\textsuperscript{57,58}

Moreover, the half-life of CPPs can be prolonged via backbone stabilization by the inclusion of β- and γ-peptide residue in the CPP sequence. In addition to increasing chemical stability, backbone stabilization may also enhance physical stability.\textsuperscript{42,59,60} Furthermore, there have been a number of strategies suggested to improve the stability of CPPs, and the reader is referred to a review done by Reissmann.\textsuperscript{60}

**Toxicity and immunogenicity**

Although CPPs are generally considered safe in terms of toxicity and immunogenicity at effective concentrations,\textsuperscript{61,62} though toxic and immunoinducing effect of some CPPs have been reported. Verdurmen and Brock reviewed CPPs and drug carriers and found a variety of side effects that might have been related or unrelated to their mechanisms of internalization.\textsuperscript{63} The toxicity of CPPs depends mainly on peptide concentration, type of cargo molecule, and conjugation strategy.\textsuperscript{64} Moreover, amphiphilic CPPs are known to be more toxic than cationic ones.\textsuperscript{55}
Despite some in vitro reports on cytotoxic effects of CPPs, it is difficult to compare results from different studies. However, in one comparative study by El-Andaloussi et al on delivery efficiency and cytotoxicity of three well-known CPPs—Tat, TP10, and penetratin—were characterized using different cargoes. The result of this study demonstrated that the cytotoxicity of these peptides was cargo dependent. In addition, the extent of cytotoxicity and internalization of TP10 varied significantly depending on the position of cargo coupling within the peptide. Among the three peptides, TP10 exhibited long-term toxic effects in HeLa and CHO cells at 20 μM. Penetratin and TAT had no effect on membrane integrity, whereas TP10 induced LDH leakage (~20%) at 10 μM in HeLa cells. On the other hand, Suhorutsenko et al evaluated cytotoxic and immunogenic responses of TP10 and its chemically modified analogues—PepFect peptides (PF3, PF4, and PF6)—in monocytic leukemia and peripheral blood mononuclear cell lines. All peptides were found to be nontoxic and nonimmunogenic in vitro at concentration of 10 μM and 5 μM, and in vivo at a dose of 5 mg/kg.

Kilk et al conducted a metabolomic approach to a CPP toxicity study on five representatives of the most common CPPs: transportan (TP), penetratin, HIV TAT-derived peptide, nona-arginine (R9), and model amphiphilic peptide. The study showed that the intracellular metabolome was mostly affected by TP, followed by HIV TAT-derived peptide and model amphiphilic peptide. Only minor changes were observed with penetratin or R9 treatment. The cells recovered from 5 μM TP treatment, but no recovery was observed at higher concentrations. This metabolomic study demonstrated that TP affected cellular redox potential and depleted energy and pools of purines and pyrimidines.

In general, few in vivo studies are available on toxic and immunological responses of CPPs. One in vivo toxicity study of CPPs and ACPPs showed that systemically administered CPPs (R9) resulted in acute toxicity in mice at a dose fourfold lower than MMP-cleavable ACPPs. The study also reported that CPPs bound to the vasculature at the site of injection, then redistributed, mostly to the liver. However, ACPPs were distributed among tissue more broadly, showed prolonged circulation, were eliminated by both renal and hepatobiliary routes, were less toxic, and enabled targeting of tumor-expressing enzymes that cleaved the linker.

An in vivo study of immunological response to delivery of a p38 siRNA using the penetratin and HIV-TAT to mouse lungs was conducted by Moschos et al. The study showed that penetratin coupled with siRNA induced an innate immunoresponse, but siRNA coupled with HIV-TAT and penetratin (not complexed with siRNA) did not. Similarly, an in vitro study reported more recently by Carter et al showed that HIV-TAT, Antennapedia and TP failed to induce a significant increase in the release of the inflammatory cytokines IL8 and IL6 in epithelial cells.

The presence and origin of cargo molecules in CPP–cargo conjugates also influences toxicity and may exert a different effect from that produced by the peptide alone. Research has indicated that toxic effects may be derived from conjugation of different cargoes to CPPs. For example, penetratin alone and TAT coupled with siRNA do not cause an immunoresection, but when penetratin is conjugated to siRNA, the complex does provoke an innate immunoresponse. The conjugation of carboxyfluorescein to CPPs is known to increase CPP cytotoxicity. Cardozo et al showed that the cytotoxicity of TA45–57 and penetratin was higher for CPP conjugates with larger cargo peptides than for those unlabeled or with smaller cargo. In addition to CPP–cargo conjugates, nanomaterials used as carriers elicit an immunoresponse and cytotoxicity.

**Activable CPPs**

The lack of cell specificity of CPPs is mainly due to their electrostatic interaction. Generally, cationic CPPs bind to anionic components of the plasma membrane, and this is principally accountable for their membrane-transporting properties. To overwhelm the problem of aspecificity, researchers introduced ACPPs. ACPPs are novel targeting agents comprising of a polycationic CPP attached to a neutralizing polyanion unit via a cleavable linker (Figure 2). Adsorption and uptake of CPPs into cells are prevented until the linker is cleaved. In ACPPs, stimulus-responsive materials are exploited to stimulate the selective display of CPPs within the pathological environment of a tumor, such as lower pH caused by buildup of lactic acid or overexpression of extracellular matrix development-remodeling proteases, or may be the external application of heat or light to a disease site.

**Acid-activated CPPs**

The lower pH of the tumor microenvironment compared with normal tissue can be used as a targeting strategy. This approach has been known to improve the intracellular delivery of cargo molecules functionalized with CPPs. A number of studies involving different approaches toward acid-activated CPPs for tumor-targeted delivery have been reported. For instance, Jin et al investigated acid-activated...
peptide by amidizing TAT lysine residue amines to succinyl amides (aTAT), completely inhibiting TAT-aspecific interactions in the blood compartment. The succinyl amides in the aTAT were rapidly hydrolyzed, fully reestablishing TAT functions in acidic tumor tissue or inside cell lysosomes/endosomes. As such, aTAT-functionalized PEG-block-poly(ɛ-caprolactone) micelles result in long circulation in the blood compartment and efficiently accumulate and transport doxorubicin (Dox) to tumor tissue, offering high antitumor activity and low toxicity to the cardiovascular system.

Cheng et al also used ACPPs with 2,3-dimethylmaleic anhydride (DMA) as a shielding group functionalized with the anticancer drug Dox to produce a novel prodrug (Dox-ACPP-DMA) for tumor-targeting drug delivery. DMA (shielding group) conjugated to ACPPs through an amide bond between DMA and the primary amines of K$_6$, which was used to inhibit the cell-penetrating activity of the polycationic CPPs (R$_8$) by intramolecular electrostatic attraction at physiological pH 7.4. However, at tumor extracellular pH 6.8, the removal of the shielding group by hydrolysis resulted in charge reversal, activating and restoring the original function of CPPs for better cellular uptake by tumor cells. After cell internalization, drug release in cells was further triggered by overexpressed intracellular proteases. In another example, Regberg et al demonstrated that peptides (PF3) modified with a leucine–histidine sequence was found to be a pH responsive CPP. The modified PF3 analogue showed significantly improved cellular bioactivity over the unmodified PepFect. Similarly, a tumor-specific pH-responsive peptide H$_2$K(R$_2$)$_7$-(RRK[HHH][HHH][HHH][RR]) has been reported to have the ability to respond to acidic pH in the tumor microenvironment.

TH(AGYLLGHINLHHLAHL[Aib]HHIL-NH$_2$) is another promising pH-responsive CPP derived by complete replacement of all lysines in the TK(AGYLLGKINLKKL[Aib]LLIL-NH$_2$) sequence of by histidines. TH does not show cell-penetrating efficacy in normal tissue or during blood circulation because of the neutral pH under those environments. However, at lower tumor environment pH, the cell-penetrating capacity of TH is activated as a result of histidine protonation in TH peptides and conversion of the surface charge of TH from negative to positive. Paclitaxel (Ptx)-loaded TH-modified liposomes show significantly greater inhibition against tumor cell growth than that of PEG liposomes and free Ptx both in vivo and in vitro at pH of 6.3 as compared with at pH 7.4.

Similarly, Yao et al developed a modified form of TH by the introduction of electron-donating group, such as ethyl, isopropyl, and butyl to the C$_2$ position of histidine TH to form corresponding TH analogues (ethyl-TH, isopropyl-TH, and butyl-TH). The TH analogues formed were conjugated to a camptothecin (Cpt)- and butyl-TH-modified conjugate and

![Figure 2: Activable cell-penetrating peptides (CPPs). Note: Enzyme-activated CPP (A), photon-activated CPP (B), pH-activated CPP (C).](image-url)
displayed a pronounced cytotoxicity effect on cancer cells more strongly in a pH-dependent manner compared to TH and other conjugates.\textsuperscript{82}

Very recently, new pH-controllable CPPs (PCCPPs) were reported on by Lee et al.\textsuperscript{83} According to this study, synthesized poly-l-lysine-based PCCPPs were capable of undergoing pH-dependent conformational transition, thereby displaying specific cell-penetrating properties at the target site. At a physiological pH, the PCCPPs contained a low helical tendency due to electrostatic attractions between carboxylate and protonated amine groups in each side chain. Therefore, unselective cellular internalization was hindered. In contrast with an acidic environment (pH 5–6), the pH-inactivated motif quickly changed to an intact helical structure, whereby electrostatic interactions between repulsions and attractions were well balanced throughout the side chains. Accordingly, the conformation of PCCP was able change to a helix that possessed greater cell-penetrating properties specifically at cancer sites.

**Enzyme-activated CPPs**

Specific enzymes, such as proteases, glycosidases, or esterases, are overexpressed in tumor cells, and this fact can be used as a tool for developing tumor-targeting drug delivery.\textsuperscript{84} Quenching of the cell-penetrating activity of polycationic peptides by electrostatic interactions with the polyanionic domain can block cellular uptake. However, in tumor tissues, MMP2/9 are overexpressed, cleave the substrate, and release polycationic peptides from the polyanionic domain, thereby stimulating cellular adhesion and subsequent uptake of peptides (Figure 2).\textsuperscript{76,85} Shi et al developed a conjugate of ACPPs with Dox, which is sensitive to MMP2/9, to form ACPP–Dox conjugates consisting of the polycationic domain (CPPs), cleavable MMP2/9-sensitive substrate, the polyanionic domain, and Dox. Activation of ACPP–Dox was found to occur via MMP2/9 in an enzyme-concentration-dependent manner. Flow cytometry and laser confocal microscopy demonstrated that higher cellular uptake of ACPP–Dox occurred with HT1080 cells (overexpressed MMPs) than MCF7 cells (underexpressed MMPs) after enzyme-triggered activation.\textsuperscript{86}

Similarly, Gao et al developed NPs functionalized with ACPPs: EEEEEEEE(E\textsubscript{3})-6-aminohexanoyl-PLGLAG-RRRRRRRR(R\textsubscript{3}) to deliver cargoes effectively to glioma cells and further improve glioma treatment. In the blood, cationic R\textsubscript{3} is shielded by E\textsubscript{3} via electrostatic forces, and the cell-penetrating capability of R\textsubscript{3} is blocked. In the glioma site, PLGLAG is cleaved by MMP2. Therefore, E\textsubscript{3} was able to be detached from R\textsubscript{3} at the glioma site, leading to a regaining of penetration ability of R\textsubscript{3}.\textsuperscript{87}

Xia et al also constructed MMP-based activable low-molecular weight protamine (ALMWP; CVSRRRRRRGGRRRR)-functionalized NPs. Ptx-loaded ALMWP NPs demonstrated superior antitumor efficacy over unmodified NPs and LMWP-functionalized NPs.\textsuperscript{88} Other research has also found that MMP2-based ACPP nanocarriers show a greater cellular uptake, cytotoxicity, tumor targeting, and antitumor efficacy in vitro and in vivo than non-MMP 2-based and free drugs.\textsuperscript{89}

Cancer cells produce high levels of intracellular reactive oxygen species (ROS), such as O\textsubscript{2}–, H\textsubscript{2}O\textsubscript{2}, and HO\textsuperscript{•}, compared to the normal ROS environment. Using this principle very recently, Yoo et al developed novel protease ACPPs containing an ROS-responsive methionine, a cell-permeable lysine chain (poly[l-methionine-block-l-lysine]) (ML), and a MMP-cleavable linker (PLGLAG). The ML–PLGLAG conjugate (MLMP) and Dox were incorporated into the micelle core. MLMP demonstrated MMP-sensitive cleavage and ROS-induced Dox release, while MLMP (Dox) exhibited pronounced tumor inhibition efficiency with no toxicity compared to free Dox. The findings of this study suggesting that dual-stimuli-based delivery systems have great potential in cancer therapy.\textsuperscript{90}

Although MMP-based ACPPs are known to target tumors effectively in vivo, ACPP substrates are not perfectly selective for MMPs, and MMP is not exclusively expressed in tumors. Therefore, in order to improve the targeting potential of ACPPs, Whitney et al explored other classes of proteases and substrates that might offer better specificity and flexibility than MMP-based approaches by using phage display. Accordingly, a unique ACPP (RLQLKL) was identified. This ACPP was not cleaved by MMP or various coagulation factors but was efficiently cleaved by elastases and plasmin, both of which have been shown to be overexpressed in tumor cells.\textsuperscript{91} Similarly, cathepsin B enzyme-responsive\textsuperscript{92} and urokinase-type plasminogen activator-based\textsuperscript{93} ACPPs have been reported for their potential in tumor-specific drug delivery systems.

**Photon-sensitive CPPs**

The two aforementioned ACPP types involve the activation of CPPs in vivo in tumor tissue. However, another approach is the external illumination of tumors by near-infrared or ultraviolet (UV) light to stimulate dissociation of photosensitive groups from CPP NPs, thus controlling the release of drugs at the tumor site.\textsuperscript{94} Shamay et al developed photon-sensitive CPPs using polymers bearing light-activated caged CPPs
for selective cellular uptake upon UV-light illumination (365 nm), which may offer a promising approach to the delivery of payloads to target cells. Lin et al also employed this strategy to take advantage of both photosensitive peptides (PSPs) and asparagine–glycine–arginine (NGR) to develop an siRNA delivery system for the treatment of CD13-overexpressing exterior tumors with the aid of UV illumination. The PSPs consisted of two units: CPPs (CKRRMKWKK) and a photolabile-protective group (Nvoc). Nvoc masked the positive charges of lysine residues on CPPs temporarily. In circulation, the penetration capability of the CPPs was shielded, but upon reaching the tumor environment, the uncaging of the PSPs was stimulated by the external application of UV light, resulting in activated CPPs. However, the use of UV light has been limited, due to its harmful effect on tissue and low penetrability. Near-IR light is able to penetrate tissues deeply and is less harmful to cells in contrast to UV light. With this in mind, Yang et al designed NLCs conjugated with photon-sensitive CPPs (psCPPs; CGR-RMKWKK) and NGR in an attempt to improve targeted delivery of Ptx to tumor cells. The psCPP unit facilitated specific cellular uptake after cleavage of the photon-sensitive protective group, whereas NGR moiety discriminately bound to CD13-positive tumors. The results of the study demonstrated that the tumor-growth-inhibition rate and cellular uptake of the psCPP/NGR-NLC group were significantly higher than the rest of the Ptx groups. Similarly, Yang et al developed near-IR light-activated psCPP/NGR liposomes that provide a reasonable approach to selective targeted delivery of siRNA.

**Thermally activated CPPs**

In this approach, a biopolymer elastin-like polypeptide, a heat-sensitive carrier able to undergo phase transition upon reaching an externally heated tumor environment is used. This carrier is tumor specific, because it aggregates only at heated tumor sites at 39°C–42°C. Such peptides as CREKA, CLT1 (CGLIIQKNEC), and CLT2 (CNAGESSKNC) are also known to target tumor blood vessels and represent as a novel type of homing peptide. They have an affinity to bind with clotted plasma proteins that exist on the walls of tumor vasculature as well as in tumor stroma. Li et al identified TCP1 (CTSPFSHC), a vasculature-homing peptide, by in vivo phage library screening in an orthotopic colorectal cancer model. The TCP1 peptide was found to recognize selectively the vasculature of orthotopic colorectal cancer in normal BALB/c mice induced by syngeneic colon cancer cells (colon 26). Recently, a number of more peptides homing to tumor vasculature have been identified.

**Tumor-targeting peptides**

TTPs are also known as tumor homing peptides and are small peptides shorter than CPPs by three to ten residues that have strong affinity and specificity for tissue targets or tumor cells. The overexpression of specific receptors in most tumors and their vasculature is utilized by TTPs, for which they show high binding ability. Among a number of different receptors that are known to be overexpressed in tumor cells, integrins are the most attractive target for drug delivery, because they have crucial roles in the process of tumor cell proliferation, migration, invasion, and survival.

**Peptides-targeting tumor vasculature**

As mentioned earlier, tumor vasculature varies from normal blood vessels both structurally and morphologically. For example, the tumor blood vessels are leaky and porous, unlike normal vasculature. In addition to the altered morphology, tumor vasculature differs from that of normal by its molecular composition. Arginine–glycine–aspartic acid (RGD) and NGR peptides are the most extensively studied peptides for targeting tumor vasculature. RGD peptides can selectively target tumor vasculature expressing α5β3, and αvβ3 integrins, and an NGR peptide (CNGRC) binds to CD13 (aminopeptidase N), which is expressed in the tumor vasculature specifically. RGR is another peptide that was selected from phage display in pancreatic tumors that have exhibited superior affinity to angiogenic vessels in insulinomas and recognize various αβ integrins. RGR has been exploited as a carrier for delivery of therapeutic proteins (TNFα and IFNγ) to targeted sites in cancer therapy. Another peptide that is targeting to angiogenic vasculature is F3. F3 (KDEPQRRSARLSAKPAPPKPEPKPKKAP AKK) is a 31-ami acid peptide that is able to target blood vessels and tumor tissue. F3 is known to target nucleolin, which is selectively expressed on the surface of endothelial cells and tumor cells. After binding, F3 is internalized through receptor-mediated endocytosis, and then transllocated from the cytoplasm to the nucleus, where it distributes itself throughout organelles.

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**Peptides-targeting tumor lymphatics**

LyP1 (CGNKTRGEC) and LyP2 (CNRTKAGC) are peptides that home to tumor lymphatics. Laakkonen et al identified LyP1 (CGNKTRGEC) by screening breast carcinoma xenografts (MDA-MB435). LyP1 binds selectively to p52, which is a mitochondrial protein that shows unusual
expression on surface tumor lymphatics, tumor cells, and a subset of myeloid cells. LyP1 has been found to distinguish lymphatics and tumor cells in MDA-MD435 and MMTV-PyMT breast carcinoma and KRB osteosarcoma xenografts and their metastatic lesions; however, it is unable to recognize C8161 melanomas. Simultaneously, the LyP2 peptide (CNRRTKAGC) homes to lymphatics of C8161 melanomas and cervical carcinomas and K14-HPV16 skin, but not to MDA-MB435 tumors, showing heterogeneity in the molecular markers of tumor cells and lymphatics. Another peptide, RMS-II (CMGNKRSAKRPC), which has some sequence similarity with LyP1, was also identified in an in vitro screen for peptides binding to RMS cell lines. Better targeting ability to RMS xenografts was observed by RMS-II than LyP1 in vivo. Furthermore, RMS-II recognized tumor blood vessels, but not tumor lymphatic vessels, which showed the different specificities of these two peptides.

Peptides homing to tumor cells

Some homing peptides are known to possess cell-penetrating properties. For example, F3 and LyP1 peptides are cell-type-specific CPPs. These peptides have the ability to enter tumor cells and blood (F3) or lymphatic endothelial cells (LyP1) in tumors to which they home. CGNKRTR (tLyP1) is also a homing and tumor-penetrating peptide via NRP1-mediated endocytosis via the C-end rule internalization pathway.

CRGRRST (RGR) and CGKRK are other homing peptides that are conveyed to a nucleus of the targeted cell after cellular uptake. These peptides consist of a number of basic amino acids, which are considered to be accountable for intracellular and intranuclear transportation. He et al synthesized a novel tumor-homing peptide, CSNIDARAC (4R22), which bound strongly to H460 cells and efficiently internalized into the cells. It also selectively bound to other lung tumor cell lines compared to cells of different cancer types.

A number of endothelial cell-targeting peptides have been found to act as tumour-penetrating peptides to enable the internalization of a conjugated drug to the cancer cell. These peptides share a specific C-terminal C-end rule sequence, (R/K)XX(R/K), which is responsible for tissue penetration and cell internalization. For example, internalizing RGD (iRGD; CRGDKGPDC), one of the most innovative TTPs is a 9-amino acid cyclic peptide, where the lysine residue can also be an arginine and the aspartic acid a glutamic acid. In addition to targeting the \( \alpha_\beta_3 \) integrin receptor, it is able to penetrate tumors. Compared to other RGD peptides, iRGD can distribute widely in extravascular tumor tissue.

Synergistic use of CPPs with TTPs in tumor-targeting delivery

TTPs bind with receptors that are upregulated on tumor cells selectively but may not be capable of reaching the target by themselves. On the other hand, CPPs penetrate plasma membrane effectively but lack target specificity. Target-specific CPPs able to both penetrate the plasma membrane and selectively transport the drug to the desired target of action is considered an ideal type of CPP.

Combining a TTP with a suitable CPP is known to facilitate the translocation of the conjugate moieties to target tumor sites with better selectivity and specificity. Studies have shown that TTPs conjugated with CPPs have dramatically increased efficiency in translocating drug molecules specifically to cancer cells compared to TTPs alone. Myrberg et al used the cyclic peptide cCPGEGAGC (PEGI), which is a homing peptide that has been known to target breast tumor tissue in mice; however, the PEGI peptide was not able to cross the plasma membrane. The combined use of PEGI with PVEC by conjugation displayed specific targeting to breast cancer cells in vitro and in vivo. In addition, the conjugated PEGA–pVEC chimeric peptide was found to increase the efficacy of chlorambucil more than fourfold. A further study by the same group used the combination of CREKA with pVEC as a chimeric peptide for delivery of chlorambucil intracellularly. The results revealed that the chlorambucil–CREKA–pVEC conjugate produced significantly enhanced anticancer activity in vitro over the anticancer drug alone. The study also showed that CREKA–pVEC was better in translocating cargo molecules inside cancer cells compared to the PEGA–pVEC peptide.

More recently, Fan et al evaluated the combined effect of SP90 (SMDPFLFQLLQL) with C peptide (GPGLWERQAREHSERSKKRRRESECKAA) in breast cancer homing ability. SP90 is able to bind selectively to breast cancer cells (MDA-MB231), but C peptide has no cell specificity. However, the combined use of SP90 with C-peptide (SP90–C) showed a 10- and 12-fold increase in efficiency of intracellular delivery compared to C peptide and SP90 alone, respectively. Furthermore, SP90 and SP90–C conjugate were utilized for the delivery of HIV1 VPR (potential novel anticancer protein drug) into breast cancer cells. SP90–VPR–C has demonstrated improved apoptosis-inducing and antiproliferative activity of HIV1 VPR, without affecting normal breast cells.

Nowadays, sequences of peptide exhibiting both cell-penetrating and specific targeting properties are designed using mRNA-display technology. For instance, RLW...
(RLWMRWYSPRTRAYG) is known to exhibit both properties and can target A549 cells specifically. Gao et al conjugated RLW with NPs to form RNPs for targeted delivery into lung cancer cells (A549), and a conventional CPP (R₈ [RRRRRRRR]) was used as a control. In vitro cell uptake study showed that RNPs specifically improved uptake by A549 cells compared to human umbilical vein endothelial cells. However, conjugation with R₈ increased uptake by both cell types, indicating better targeting specificity for RNPs. RNPs loaded with docetaxel (Dtx) demonstrated RLW selectively targeted A549 cells and enhanced the cytotoxicity of the drug in vitro.¹¹ In addition, Lim et al developed a novel nontoxic and cancer-specific CPP, BR2 (BR2RAGLQFPVGRLLRLLR [17 amino acids]), based on the cell-penetrating motif of buforin IIIB. BR2 was fused to a single-chain variable fragment (scFv) antibody against mutated Kras. BR2 exhibited cancer-specific activity without toxicity to normal cells. Moreover, BR2 displayed greater membrane translocation efficiency than TAT.⁴⁹⁻⁵⁷ Similarly, BR2 fused with scFv was found to induce a higher degree of apoptosis than TAT-fused scFv in Kras-mutated HCT116 cells.¹¹⁰

**Peptides for targeting MDR cancer**

Most recently, anticancer drug–CPP/TTP conjugates have been one of the promising strategies to overcome tumor MDR. Sheng et al designed a dual-targeting hybrid peptide – HAIYPRHGCGMPKKKPTPIQLNP (T10-ERK) – composed of an ERK-peptide inhibitor (MPKKKPTPIQLNP), a thiol linker (GGCG), and transferrin receptor-binding peptide (HAIYPRH) to overwhelm the problem of drug resistance. T10-ERK was conjugated to DoxO-EMCH (a prodrug of Dox), resulting T10-ERK–Dox. The efficacy of T10-ERK–Dox in reversing drug resistance compared with free Dox, and T10–Dox was determined using MCF7/ADR cells and nude mice-bearing MCF7/ADR xenografts. The results of the study indicated that T10-ERK–Dox efficiently inhibited drug resistance and improve the cytotoxicity of Dox by blocking Pgp-mediated drug efflux and inducing apoptosis.¹¹²

Furthermore, Lelle et al formulated a conjugate consisting of octaarginine CPP (Ac-CRRRRRRR-NH2) and Dox dimer with high DNA affinity. The cytotoxicity of the peptide-drug conjugate was evaluated against drug-sensitive and Dox-resistant cancer cells, and showed that the conjugate overcame drug resistance in neuroblastoma cells efficiently.¹¹¹ Feng et al also studied a Dox–SAPSP (slightly acidic pH-sensitive peptide) conjugate in order to interfere with drug resistance in cancer therapy. In this study, Dox was attached to SAPSP (CHGAHEHAGHEHHAHE-NH₂) to achieve a Dox–SAPSP prodrug. Cell uptake studies showed that Dox–SAPSP selectively accumulated in both Dox-sensitive and Dox-resistant cancer cells and was 26-fold less toxic toward noncancerous MCF10A cells than free Dox.¹²²

**Application of CPP/TTP nanocarriers for herb-based anticancer drugs**

**Herb-based anticancer drugs**

The use of plants or phytoconstituents has a long history in cancer therapy and has played a significant role in the discovery of effective anticancer agents. Approximately 60% of presently used anticancer agents are derived from natural products in one way or another.¹²³ Recently, more than 3,000 plants globally were reported to have anticancer properties.¹²⁴ Some examples of important anticancer drugs that are derived from plant sources including Ptx (Taxol), vinblastine, capsaicin, vincristine, the Cpt derivatives, topotecan, irinotecan, and etoposide (Table 1 and Figure 3).¹²⁵

Vinca alkaloids (vinblastine and vincristine) are obtained from the Madagascar periwinkle, Catharanthus roseus (Apocynaceae), and were the first plant source used clinically as anticancer agents for leukemias, lymphomas, breast, testicular, and lung cancers, and Kaposi’s sarcoma.¹²⁶ In recent years, semisynthetic derivatives of vinca alkaloids, such as vindesine and vinorelbine, have received approval from the US Food and Drug Administration, and vinflunine has been approved by the European Medicines Agency as a second-line chemotherapeutic agent in the treatment of metastatic urothelial cancer.¹²⁷ Moreover, vinflunine and vinorelbine have shown minimized toxicity in comparative animal models.¹²⁸,¹²⁹

Taxanes are a class of herbal drugs that are commonly utilized in the treatment of breast cancer and initially isolated from the plant (Taxus brevifolia). Ptx (Taxol) and docetaxel (Taxotere) are the derivatives of taxanes that are clinically very essential in combination with synthetic chemotherapeutic agents such as Dox, capetcitabine, and gemcitabine in order to achieve better clinical effects.¹³⁰ Cpt is a quinoline alkaloid with topoisomerase I-inhibitor activity obtained from the bark and stem of Camptotheca acuminata.¹³¹ However, it was withdrawn from clinical trials due to its poor water solubility and severe toxic effects. In order to solve these limitations of Cpt, a number of Cpt analogues, such as topotecan, irinotecan, and belotecan, have been derived and approved for clinical use.¹³²
Table 1  Plant-derived anticancer agents in clinical use and clinical trials

<table>
<thead>
<tr>
<th>Anticancer agent</th>
<th>Source</th>
<th>Chemical class</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Podophyllotoxin</td>
<td>Podophyllum peltatum, P. emodi</td>
<td>Podophyllotoxin</td>
<td>123</td>
</tr>
<tr>
<td>Etoposide</td>
<td>P. peltatum</td>
<td>Podophyllotoxin</td>
<td>123</td>
</tr>
<tr>
<td>Teniposide</td>
<td>P. peltatum</td>
<td>Podophyllotoxin</td>
<td>123</td>
</tr>
<tr>
<td>Combretastatin A4 phosphate</td>
<td>Combretum caffrum</td>
<td>Stilbenes</td>
<td>126</td>
</tr>
<tr>
<td>Betulinic acid</td>
<td>Betula spp.</td>
<td>Triterpene</td>
<td>126</td>
</tr>
<tr>
<td>Topotecan</td>
<td>Camptotheca acuminata</td>
<td>Alkaloid</td>
<td>127</td>
</tr>
<tr>
<td>Irinotecan</td>
<td>Plants of the genus Taxus (yews)</td>
<td>Alkaloid</td>
<td>127</td>
</tr>
<tr>
<td>Docetaxel</td>
<td>Plants of the genus Taxus</td>
<td>Taxane</td>
<td>127</td>
</tr>
<tr>
<td>Camptothecin</td>
<td>Camptotheca acuminata</td>
<td>Quinoline alkaloid</td>
<td>127</td>
</tr>
<tr>
<td>Flavopiridol</td>
<td>Dysoxylum binestariferum</td>
<td>Flavonoid alkaloid</td>
<td>127</td>
</tr>
<tr>
<td>Curcumin</td>
<td>Curcuma longa</td>
<td>Polyphenol</td>
<td>127</td>
</tr>
<tr>
<td>Elliptinium</td>
<td>Bleekeria vitensis</td>
<td>Alkaloid</td>
<td>132</td>
</tr>
<tr>
<td>Gambogic acid</td>
<td>Garcinia hanburyi</td>
<td>Xanthohoid</td>
<td>136</td>
</tr>
<tr>
<td>Ingenol mebutate</td>
<td>Euphorbia peplus</td>
<td>Diterpene</td>
<td>137</td>
</tr>
<tr>
<td>Homoharringtonine</td>
<td>Cephalotaxus harringtonia</td>
<td>Alkaloid</td>
<td>137</td>
</tr>
<tr>
<td>Vinblastine</td>
<td>Catharanthus roseus</td>
<td>Vinca alkaloids</td>
<td>137</td>
</tr>
<tr>
<td>Vincristine</td>
<td>C. roseus</td>
<td>Vinca alkaloids</td>
<td>137</td>
</tr>
<tr>
<td>Paclitaxel</td>
<td>Taxus brevifolia</td>
<td>Taxane</td>
<td>137</td>
</tr>
<tr>
<td>Genistein</td>
<td>Genista tinctoria</td>
<td>Flavonoid</td>
<td>138</td>
</tr>
<tr>
<td>Resveratrol</td>
<td>Polygonum cuspidatum</td>
<td>Flavonoid</td>
<td>138</td>
</tr>
<tr>
<td>Piperlongumine</td>
<td>Piper longum</td>
<td>Alkaloid</td>
<td>139, 140</td>
</tr>
<tr>
<td>Brucelantin</td>
<td>Brucea antisymesiterica</td>
<td>Quassinoid</td>
<td>141</td>
</tr>
</tbody>
</table>

Podophyllotoxin is another essential active antitumor agent extracted from the roots and rhizomes of Podophyllum spp. (Berberidaceae). The two semisynthetic derivatives of podophyllotoxin that are used in the treatment of lymphomas and bronchial and testicular cancers are etoposide and teniposide. Homoharringtonine and elliptinium are the other herb-derived agents in clinical use. Homoharringtonine was originally derived from the Chinese plant Cephalotaxus harringtonia var. drupacea (Cephalotaxaceae) and exhibits excellent anticancer activity against different types of leukemias, including some resistant to standard treatment. A racemic mixture of harringtonine and homoharringtonine is being utilized effectively in China to treat both acute and chronic myelogenous leukemia. Elliptinium was isolated from species of many genera of the family Apocynaceae, including Bleekeria vitensis, a Fijian medicinal plant with anticancer properties.

There are a number of promising anticancer drugs currently in clinical trials. For instance, Piperlongumine is a natural product extracted from Piper longum (Piperaceae) known to have potent anticancer activity. Curcumin is a polyphenol extracted from Curcuma longa. Despite curcumin’s poor absorption, a Phase II trial has shown anticancer activity in some patients with pancreatic cancer, and it is known to be well tolerated. Among bioactive compounds from medieval European plants, resveratrol, garlic compounds, and hypericin are under clinical evaluation. Besides these, a large number of other bioactive compounds isolated from herbs are under clinical investigation. In addition, several bioactive agents isolated from Chinese herbs have recently been found to exhibit strong anticancer activities, and are undergoing preclinical and clinical studies. Among these natural products with anticancer potential are flavonoids (gambogic acid, curcumin, silibinin, and wogonin), alkaloids (berberine), terpenes (artemisinin, β-elemene, ursolic acid, triptolide, and oridonin), quinones (emodin and shikonin), and saponins (ginsenoside Rg3).

Recent examples of herb-based anticancer-bioactive CPP/TTP nanocarriers

Though a large number of potent plant-derived anticancer drugs have been investigated, their clinical use is limited, due to the fact that most herbal bioactive extracts are hydrophobic in nature, resulting in poor bioavailability and therapeutic failure. To circumvent this challenge, different nanocarriers, such as NPs, nanocapsules, liposomes, quantum dots, phytosomes, dendrimers, and nanoemulsions, have been exploited to deliver herbal anticancer bioactive agents. As mentioned in the previous sections, nanochemotherapeutic agents exhibit improved bioavailability, enhanced bioactivity, and stability with minimal systemic toxic effects.

Moreover, surface modification of these nanocarriers with various ligands has been found to improve the targeting...
of bioactive agents to a cancer cell and then increase therapeutic efficacy and reduce toxic effects. Among these, CPPs/TTPs provide a highly promising approach for the intracellular delivery of anticancer drugs. In this section, we discuss the delivery of herb-based anticancer drugs using CPP/TTP-modified nanocarriers. Several studies have reported that CPP/TTP–herb-based anticancer conjugates demonstrate promising antitumor activity (Table 2).

Curcumin is an extensively investigated active flavonoid isolated from the rhizome of *C. longa* with broad-spectrum anticancer properties. However, its low water solubility and poor bioavailability have limited its clinical use. In the last few decades, it has been studied in various delivery systems.
Table 2 Examples of plant-derived anticancer drugs conjugated with peptides

<table>
<thead>
<tr>
<th>Anticancer agent</th>
<th>Peptides</th>
<th>Nanocarrier</th>
<th>Application</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tripterine</td>
<td>Ste-R_L2</td>
<td>Nanostructured lipids</td>
<td>PC3</td>
<td>81</td>
</tr>
<tr>
<td>Curcumin</td>
<td>GHHNGR</td>
<td>Nanoliposomes</td>
<td>MCF7, MDA-MB468</td>
<td>144</td>
</tr>
<tr>
<td>Trichosanthin</td>
<td>HBD (GPGLWERQAREHSERKRRRESECKAA)</td>
<td>Liposomes</td>
<td>HeLa cells, 95D, A549, SMMC7721, MCF7</td>
<td>149</td>
</tr>
<tr>
<td>Paclitaxel</td>
<td>TAT (Cys-AYGRKKRRQRRR)</td>
<td>Polymersomes</td>
<td>B16F1</td>
<td>151</td>
</tr>
<tr>
<td></td>
<td>iRGD</td>
<td>Liposomes</td>
<td>B16F1</td>
<td>155</td>
</tr>
<tr>
<td></td>
<td>iRGD</td>
<td>Liposomes</td>
<td>B16F1</td>
<td>154</td>
</tr>
<tr>
<td></td>
<td>c(RGDfK)</td>
<td>Lipid–polymer hybrid</td>
<td>A2780/Pcx</td>
<td>158</td>
</tr>
<tr>
<td>Docetaxel</td>
<td>RIPL (IPLVVPRLRRRRRRRC)</td>
<td>Liposome</td>
<td>SKOV3, MCF7, DU145, PC3</td>
<td>157</td>
</tr>
<tr>
<td>Camptothecin</td>
<td>RGD-mimetic</td>
<td>Lipid–polymer hybrid NPs</td>
<td>C6 glioma cells</td>
<td>160</td>
</tr>
<tr>
<td>10-Hydroxycamptothecin</td>
<td>c(RGDyK)</td>
<td>NPs</td>
<td>MDA-MB231</td>
<td>161</td>
</tr>
</tbody>
</table>

Abbreviations: PEG, polyethylene glycol; PLA, polyactic acid; NPs, nanoparticles; RGD, arginine–glycine–aspartic acid.

to circumvent these limitations. Kangarlou et al synthesized linear tumor-homing peptides (GHHNGR) and conjugated them with curcumin-loaded nanoliposomes. The conjugated curcumin-loaded liposomes showed significant cytotoxicity around sevenfold that of an aqueous curcumin suspension in MCF7 (IC50 3.8 μM) and MDA-MB468 (IC50 5.4 μM). Furthermore, the entrapped curcumin exhibited a prolonged half-life and reduced degradation, in contrast to the free drug, in aqueous suspensions. Simion et al also demonstrated that lipid nanoemulsion-loaded curcumin (CmLN) conjugated with a nona-arginine peptide (R9–CmLN) exhibited significantly higher uptake and internalization of R9–CmLN compared to nonfunctionalized CmLN in human endothelial cells. Furthermore, Das et al synthesized RGDK–lipopeptides and functionalized these noncovalently with single-walled carbon nanotubes (SWNTs) to form RGDK–SWNT. RGDK–SWNT was capable of delivering the anticancer drug curcumin to B16F10 melanoma cells more efficiently than NIH3T3 cells (noncancerous), leading to the selective killing of B16F10 cells.

Tripterine, also known as celastrol, is a Chinese herbal medicine extracted from the thunder-god vine (Tripterygium wilfordii) and demonstrates promising bioactivity against a number of tumor cell lines. Yuan et al studied the antitumor activity of tripterine using CPP (Ste-R_L2)-modified NLCs in a human prostate cancer cell line (PC3) and a prostate tumor-bearing mouse model. CPP–tripterine (CT) NLCs showed dramatically enhanced antitumor activity in vivo and in vitro in comparison with free tripterine. In vivo tumor inhibition rates of CT-NLCs (54.5%) were higher than cyclophosphamide (37.07%) at low doses (2 mg/kg) in the PC3 cell line. Moreover, significantly higher amounts of TNFα and IL6 cytokine content were observed following the administration of CT-NLCs and T-NLCs than high-dose tripterine.

Trichosanthin (Tcs) is a protein extracted from the root tubers of Trichosanthes kirilowii and has been used as an abortifacient for centuries in China. It is a type I RIP and well-known traditional Chinese medicine for various types of tumor cells. Lu et al used a human-derived CPP (HBD) (GPGLWERQAREHSERKRRRESECKAA) to improve the delivery of Tcs. In this study, HBD was fused with the C terminus of recombinant Tcs (rTcs) to improve the translocation efficiency of Tcs. The IC50 of rTcs-HBD in the tested tumor cells was much lower than rTcs, showing that HBD delivered the rTcs into tumor cells efficiently.

Dual-modified nanocarriers with more than one ligand are gaining much attention in anticancer drug research. Chen et al developed bifunctional NPs (BF-NPs) based on PLGA–PEG and modified them with CPP (R-) and folic acid simultaneously. The vincristine sulfate-loaded BF-NPs were prepared by an emulsion solvent evaporation method. Higher cellular uptake was found for BF-NPs than NPs modified by folic acid or R-, alone. In vitro cytotoxicity, cell apoptosis, and cell cycle also exhibited better potency of BF-NPs compared to those NPs merely modified by folic acid or R-.

Ptx is a bioactive agent that has attracted much attention over the last three decades. It is a member of the taxane family, and one of the most important and effective antineoplastic agents for the treatment of many forms of advanced and refractory cancers. Ptx is a unique chemotherapeutic agent that is available on the market in three nanoplatforms—liposomes (Lipusu), polymeric NPs (Abraxane), and polymeric micelles (Genexol, Nanoxel, and Palcical) — for parenteral administration.
Several reports have demonstrated that Ptx conjugates with different types of CCP/TTP show better anticancer activity. Fu et al used the well-known CPP TAT (Cys–AYGRKKRRQRRR) and functionalized it with Ptx-loaded liposomes containing PEG for prolonged circulation. PEG was detached at the tumor site in the presence of an exogenous reducing agent (glutathione), and this allowed cell internalization of TAT. Ptx–TAT liposomes with glutathione demonstrated higher cellular uptake and better penetration to three-dimensional tumor spheroids in vitro compared with analogous stable, shielded liposomes. Moreover, improved tumor distribution and superior delivery efficiency were exhibited by Ptx–TAT liposomes in vivo. Simón-Gracia et al developed Ptx-loaded iRGD polymersomes that showed enhanced peritoneal tumor growth inhibition and suppression of local dissemination effects, in contrast to Ptx polymersomes or Abraxane. Duan et al also formulated Ptx-CPPs using TAT and the LMWP VSRRRRRRGGGR-R. Ptx-CPPs showed considerably enhanced cellular uptake, high cytotoxicity, elevated cell apoptosis, and minimized mitochondrial membrane potential against both A549 and A549T cells compared with free Ptx. Ptx–LMWP displayed stronger tumor growth inhibition than Ptx–TAT in A549T cells. There have been a number of more recent reports on the role of CPPs in the delivery of Ptx, presented in Table 2.

Docetaxel (Dtx) is a semisynthetic derivative of Ptx that is considered one of the most promising antitumor drugs and is effective against a wide range of cancer, such as prostate, breast, pancreatic, lung, gastric, and hepatic carcinomas. Recently, Yoon et al designed an RIPL peptide (IPLVVLRRRRRRRC) and conjugated it with liposomes loaded with Dtx (Dtx–RIPL) in order to deliver the drug into hepsin-expressing cancer cells selectively. Dtx–RIPL liposomes were able to suppress tumor growth and elongate survival time significantly in BALB/c nude mice with ovarian tumors (SKOV3 cells).

**Conclusion and future perspectives**

In the treatment of cancer, drug targeting is indispensable in the selective and quantitative accumulation of a drug in the target organ or tissue. NP-based targeting is of core importance in both passive- and active-targeting systems. NPs have been found to have enormous advantages in terms of reducing aspecific cellular uptake and side effects, elongating circulation, and offering controlled release and encapsulation of multiple drugs for combination therapy.

NPs modified with CPPs/TTPs could have potentially increased cellular uptake and cytotoxicity to cancer cells. A number of studies have demonstrated that CPP-NPs are able to enhance extracelluar and intracellular internalization of various small drug molecules and biomolecules. However, lack of cell specificity and in vivo degradation were found to be the main limitations of further clinical development of CPPs. The use of ACPPs and/or the use of CPPs with TTPs in combination have been known to reduce problems associated with aspecificity of CPP. More recently, researchers have developed a CPP that exhibits both cell-penetrating properties and tumor cell-targeting efficiency using mRNA-display technology. Moreover, dual-targeting hybrid–peptide conjugates and multifunctionalized NPs have been among promising strategies to overcome tumor MDR.

Despite the promising potential of CPP/TTP-modified nanocarriers for cancer therapy, there have been no adequate reports on their in vivo toxicity or immunogenicity. In addition, the issue of detailed mechanisms of penetration and pharmacokinetics and development of a sensitive method of detection of CPPs/TTPs should be thoroughly addressed for further preclinical and clinical study. Finally, we hope that more advanced and optimized CPPs will be discovered and play a significant role in shifting the paradigm of cancer therapy.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


