Chapter 21

Circumventing Tumor Resistance to Chemotherapy by Nanotechnology

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Abstract

Patient relapse and metastasis of malignant cells is very common after standard cancer treatment with surgery, radiation, and/or chemotherapy. Chemotherapy, a cornerstone in the development of present day cancer therapy, is one of the most effective and potent strategies to treat malignant tumors. However, the resistance of cancer cells to the drugs remains a significant impediment to successful chemotherapy. An additional obstacle is the inability of chemotherapeutic drugs to selectively target tumor cells. Almost all the anticancer agents have severe side effects on normal tissues and organs. The toxicity of currently available anticancer drugs and the inefficiency of chemotherapeutic treatments, especially for advanced stages of the disease, have limited the optimization of clinical drug combinations and effective chemotherapeutic protocols. Nanomedicine allows the release of drugs by biodegradation and self-regulation of nanomaterials in vitro and in vivo. Nanotechnologies are characterized by effective drug encapsulation, controllable self-assembly, specificity and biocompatibility as a result of their own material properties. Nanotechnology has the potential to overcome current chemotherapeutic barriers in cancer treatment, because of the unique nanoscale size and distinctive bioeffects of nanomaterials. Nanotechnology may help to solve the problems associated with traditional chemotherapy and multidrug resistance.

Key words: Cancer chemotherapy, Drug resistance, Nanomedicine, Nanotechnology

1. Obstacles to Cancer Treatment and the Potential of Nanotechnology

Cancer remains one of the main causes of death in humans and thus great efforts have been undertaken to develop cancer treatments (1). Cancer cells are notorious in their resistance to chemotherapy in the clinic. In fact, an enormous body of research strongly suggests that drug-resistant cancer cells that remain alive after chemotherapy are responsible for the reappearance of tumors and the poor prognosis for patients. The occurrence of drug resistance leads to the failure of tumor treatment. This is a difficult obstacle to overcome, as tumor resistance mechanisms have various origins.
It is known that several members of the ATP-binding cassette (ABC) transporter family play an important role in cancer cell with resistance to different drugs (2). Studies have demonstrated that ABCB1, ABCC1, ABCG2, and other members of the ABC family are expressed in different types of human cancers, and their expression is related to the outcome of chemotherapy: the higher their expression in a tumor, the more resistant cancer cells are to chemotherapy. For example, a poor survival rate characterizes gliomas and tumor-derived endothelial cells that express ABCB1 and ABCC1 subfamily members (3). In the tumor tissue, tumor resistance can be connected to the physiology of the tumor tissue, including a poor vasculature and unsuitable physicochemical conditions. To overcome drug resistance, many attempts have been made using strategies that consider the different chemotherapeutic mechanisms either at the cellular level or at the tissue level.

In the clinic, multidrug resistance (MDR) occurs in over 50% of patients whose cancer relapses, accounting in large part for the high mortality associated with cancer. Tumor resistance to chemotherapy in the clinic can be due to the inefficient distribution of drug relative to its targeted tumor tissue. MDR may become evidence either as a lack of tumor size reduction or as a clinical relapse after an initial positive response to antitumor treatment (4).

In clinical practice, drug resistance constitutes the failure of a patient to achieve a complete or partial response to therapy. In the laboratory, however, drug resistance is a cellular phenomenon and reflects the inability to demonstrate cytotoxicity at physiologically achievable drug concentrations in cancer cells. Drug resistance may be considered to be either intrinsic or acquired. Frequently, resistance is intrinsic to the cancer at the beginning, but as therapy becomes more and more effective, acquired resistance also becomes common. Intrinsic resistance occurs when
tumor cells are capable of escaping exposure or repairing damage induced by the cytotoxic effects of chemotherapy at initial exposure. Finally, acquired resistance dominates when resistant cells survive from a population that was initially sensitive to chemotherapy. Both intrinsic and acquired resistance may operate along several different pathways, including decreased drug accumulation, decreased drug activation, increased repair of drug-induced damage, altered drug targets, altered gene expression and drug barriers (2). The development of resistance to chemotherapy is frequently associated with broad cross resistance even to structurally dissimilar drugs, suggesting the existence of more than one potential mechanism of resistance. Multiple changes often appear simultaneously in highly resistant tumor cell lines. This observation has led to the widely accepted hypothesis that tumor resistance to chemotherapy is usually multifactorial.

Nanotechnology has the potential to overcome current obstacles to chemotherapy, because of the unique properties of nanoparticles (1–100 nm) (5). For example, solid tumors have unique features, such as leaky tumor blood vessels and defective lymphatic drainage, that promote the delivery and retention of macromolecules or nanoscale particles, a phenomenon recognized as the enhanced permeability and retention (EPR) effect. Nanoparticles can be constructed at a certain size for effective biodistribution and accumulation in the tumor. Nanoparticles are characterized by self-assembly, stability, specificity, drug encapsulation and biocompatibility as a result of their material composition (6). Many researchers now make investigation to find out how to employ nanotechnology to overcome tumor multidrug resistance in vivo and in vitro.

A major problem limiting the success of many anticancer agents lies in their inability to target tumor cells and tissues selectively. Therefore, almost all anticancer agents result in severe side effects to normal tissues and organs. In chemotherapy, pharmacologically active concentrations of an anticancer drug in the tumor tissue are often reached at the expense of massive contamination of the rest of the body. This poor specificity creates a toxicological problem that represents a serious obstacle to effective antitumor therapy. Recently, progress has been made on the design of nanoparticles with surface properties that allow better accumulation in tumor tissue after systemic administration. To improve the specificity of nanoparticles, a molecular recognition moiety is connected to the surface of the nanoparticles to target cancer cells in tumor tissue after intravenous administration.
For example, folic acid has been used to be conjugated to surface of nanoparticles (7). The rationale behind the choice of folic acid as a targeting moiety is that folic acid binding proteins are frequently overexpressed on the surface of human cancer cells. The folate-coated nanoparticles showed a tenfold higher apparent affinity for the folate binding protein than the free folate, as measured by surface plasma resonance. This increased apparent affinity was attributed to the fact that the particles represent a multivalent form of the folic acid ligand and can therefore display stronger interactions with the folate receptor. Thus, it could be expected that the folate-grafted nanoparticles would also strongly interact with the surface of malignant cells on which the folate binding protein can be overexpressed; such binding can eventually promote endocytosis of the nanoparticles mediated by folate binding protein. Indeed, only the cancer cells overexpressing the folate binding protein showed intensive uptake of the folate-decorated nanoparticles. The cancer cells that did not express the folate binding protein on the cell surface did not show any uptake of those nanoparticles. In addition, none of the cells was able to internalize PEG-coated nanoparticles without folate coating. The development of various nanoparticles with different ligands now offers a choice for targeted tumors with drug resistance. The suspension of nanoparticles is very stable, as evaluated by size measurements, and can be lyophilized. The surface properties, including the zeta potential, complement activation and protein adsorption pattern, are defined by the nature of the materials used to synthesize nanoparticles. Indeed, the biological activity of heparin grafted on the surface of nanoparticles was preserved at a level of 70% when compared to the activity measured for a heparin solution. The variety of biomolecules that can be conjugated to nanoparticles offers many possibilities for the design of targeted nanoparticles using a biomimetic approach.

Chemotherapies should ensure a specific toxic effect against the targeted tumor cells, even if the increased complexity of the outer surface obstructs their diffusion into tumor. Utilizing a ligand that binds specifically to its receptor on a malignant cell may help to reduce the dose-limiting cytotoxicity of the drug and also enable the drug to bypass the drug resistance mechanism especially caused by P-glycoprotein (Pgp) overexpression, via internalization through receptor-mediated endocytosis. This strategy not only targets the malignant cells directly, but also aims at destroying nonmalignant tumor components that are crucial for tumor survival and development. Heparin-paclitaxel-Folic Acid with its highly specific tumor uptake and potent antitumor properties fits the profile of this strategic requirement very well. More recently, liposomes have been modified by conjugating them with monoclonal antibodies directed against tumor antigens (8). Furthermore, copolymer nanoparticles can form a shell...
with a hydrophobic inner part that contains the drug. They have a kinetic behavior similar to that of liposomes. These vectors may carry drugs, radioisotopes and/or labeling agents, and are directed against the specific surface structure of tumor cells, hence increasing the specific distribution and accumulation of drugs within tumors.

Drug resistance is known to develop through a variety of molecular mechanisms within the tumor (Fig. 21.1), and various approaches overcoming tumor resistance to chemotherapy are based on various pathways (9). For example, the enzyme glucosylceramide synthase (GCS), responsible for bioactivation of the proapoptotic mediator ceramide to a nonfunctional moiety glucosylceramide, is found to be overexpressed in many multidrug-resistant tumor types and has been implicated in cell survival in the presence of chemotherapy. In an attempt to circumvent the mechanisms that cancer cells use to avoid cell death following chemotherapy, a polymeric nanoparticle was created to deliver ceramide, which triggers resistant cells to apoptosis under paclitaxel treatment. Treatment with the multifunctional nanoparticle produced 100% mortality among cultured cells. To overcome MDR in a human ovarian cancer cell line, modified poly(epsilon-caprolactone) (PEO-PCL) nanoparticles were used to encapsulate and deliver therapeutic agents for enhanced efficacy (9). With nanoparticle drug delivery, the resistant cells can be sensitized to paclitaxel near the IC50 concentration of sensitive cells. Chemotherapy enhanced via nanoparticle delivery has a promising potential as a strategy to overcome MDR.

Tumor cells can develop simultaneous resistance to multiple anticancer drugs (2). An alternate strategy suggested for overcoming MDR is association of the drug with nanoparticles (10). The rationale behind this strategy is to increase the intracellular concentration of the drug and other agents using endocytosis. Doxorubicin, an anticancer drug widely used in cancer therapy and a known substrate of Pgp, was encapsulated in various types of nanoparticles. The sensitivity of resistant cells to the doxorubicin-loaded nanoparticles was then evaluated by measuring the cytotoxic effect produced by increasing the concentration of the doxorubicin-loaded nanoparticles. Resistant cells treated with alginate or lactide-co-glycolide modified nanoparticles showed the same sensitivity to the treatment as the free drug (11). In contrast, resistant cells treated with doxorubicin-loaded poly(alkyl cyanoacrylate) (PACA) nanoparticles showed a much higher sensitivity to the drug, relative to the free drug. The sensitivity of
the resistant cells even reached the level of sensitivity of the parent sensitive cells, suggesting that the PACA nanoparticles can totally overcome the tumor resistance to doxorubicin (12). To explain the ability of nanoparticles to overcome doxorubicin resistance, researchers have proposed a mechanism based on the adhesion of the nanoparticles to the surface of resistant cancer cells. Adhesion is followed by the simultaneous release of the drug and nanoparticle degradation products that pass through the cell membrane without being recognized by Pgp.

To circumvent MDR, some proposed the use of competitive inhibitors such as verapamil. However, the clinical use of verapamil to overcome MDR is limited due to the serious adverse effects of this compound. More recent studies that have been designed to further improve the efficacy of nanoparticles in overcoming MDR have been based on limiting the activity of Pgp. This strategy is also an interesting alternative to promote the efficacy of doxorubicin-loaded nanoparticles. Soma et al. suggested co-encapsulating doxorubicin and cyclosporin A within the nanoparticles (12). Cyclosporin A is a chemosensitizing compound that can bind to Pgp and inhibit the pump efflux activity. Doxorubicin was incorporated within the core of the nanoparticles while cyclosporin A was located at the nanoparticle surface. Using different formulations of the drug-loaded nanoparticles, it was shown that the association of both doxorubicin and cyclosporin A within a single nanoparticle led to the most effective growth rate inhibition of the resistant cells. The association of cyclosporin A with doxorubicin nanospheres would also ensure that cyclosporin A reaches the same sites with the anticancer drug and also reduces its toxic side-effects. Other strategies proposed to regulate the expression of the Pgp have involved using siRNA (13, 14). However, the results obtained were disappointing because of the long half-life of Pgp, making its down-regulation difficult (14).

4. Engineered Nanoparticles Facilitates Targeting of Tumors

Nanoparticles have the potential to enhance the protection of drugs against biotransformation and rapid clearance in vivo (15). In order to do so, they must have long-circulating properties to reach the tumor tissue. In addition, they should have the proper biodistribution to target the tumor. With these objectives, studies have focused on customization of the surface properties of nanoparticles. Researchers have sought to modify the nanoparticle biodistribution to target tumors using poly(ethylene glycol) (PEG) as a coating material at the nanoparticle surface in order to reduce protein adsorption and complement activation (16). PEG-coated
nanoparticles were prepared from a poly(PEG cyanoacrylateco-
hexadecyl cyanoacrylate) copolymer (17). These nanoparticles cir-
culated longer in the blood stream, while their uptake by the liver
was reduced (18). They were found to accumulate in the brain to
a larger extent than other formulations, including the non-PEG-
coated nanoparticles (19, 20). The concentration of PEG-coated
nanoparticles in the central nervous system was shown to be greatly
increased, especially in the white matter when compared to con-
ventional nanoparticles. Recently, these nanoparticles were shown
to accumulate specifically in a glioma implanted into a rat brain.
The accumulation was found to occur mainly in the tumoral tis-
sue, while the amount of nanoparticles found in the adjacent
healthy tissue and in the control hemisphere was much lower (21,
22). The comparable distribution in tumor and normal tissue was
attributed to the difference in the microvascular permeability
between healthy and tumor tissue, combined with an increased
circulation time in the blood stream.

Maeda et al. found that Evans blue dye, which binds with
plasma albumin, concentrated selectively in tumor tissues follow-
ing intravenous (i.v.) injection (23). The same behavior was also
noticed with radiolabeled plasma proteins, including transferrin
(90 kDa) and IgG (160 kDa), whereas smaller proteins such as
neocarzinostatin (12 kDa) did not accumulate in tumors (24).
The tumor accumulation reaches up to several fold higher than
that of the plasma due to lack of efficient lymphatic drainage in the
solid tumor; this provides an ideal application for EPR-based selec-
tive anticancer drug delivery and distribution in a tumor. Tumor
blood vessels are thought to have relatively large pore structures
and poorly aligned defective endothelial cells lacking a smooth
muscle layer (25). Extensive production of vascular permeability
enhancing factors, such as nitric oxide (NO), lead to highly abnor-
mal transport dynamics across tumor capillaries, especially for
nanosized macromolecular drugs. Thus, it becomes possible for
anticancer nanomedicines of certain sizes to cross selectively into
tumor tissues (26). Furthermore, tumor tissues usually lack effect-
ive lymphatic drainage (27, 28), which leads to prolonged reten-
tion of nanoparticles. Due to their size, nanoscale particles
containing anticancer drugs administered intravenously (i.v.) can
escape renal clearance. Often they cannot penetrate the tight
endothelial junctions of normal blood vessels, but can extravasate
in tumor vasculature and become trapped in the tumor vicinity.

Establishment of this principle hastened the development of
various multifunctional nanoparticles for targeted cancer chemo-
therapy. Indeed, this highly selective local distribution of nano-
particles in tumor tissues has proven superior in therapeutic effect
with minimal side effects in both preclinical and clinical settings.
Gabizon et al. found that 100 nm nanoparticles can passively
enter tumor tissues, thereby, increasing selectivity of anticancer
drug delivery at the tumor site, while markedly reducing drug accumulation and toxicity in many susceptible healthy tissues (29). If the level of drug resistance is comparable to the drug levels in tumor, MDR may be overcome by increasing delivery of anticancer drugs based only on mass action (30).

Biocompatible and sterically stabilized micelles (SSMs) have been used as nanocarriers for chemotherapeutic agents. Drug solubilization in SSMs is reproducible and is attributed to the avoidance of drug aggregate formation. Furthermore, SSMs composed of polyethylene glycol (PEGylated) phospholipids are attractive nanocarriers for drug delivery because they are sufficiently small (14 nm) to cross through the leaky microvasculature of tumors and penetrate tissues for passive targeting of solid cancers in vivo, resulting in high drug concentration in tumors and reduced drug toxicity to the normal tissues (31).

During the past few years, several strategies have been investigated to improve the clinical effectiveness of chemotherapy; in particular, attention has been focused on drugs and their pharmaceutical formulations. A promising strategy is the use of carriers to transport drugs that are already employed in the clinic such as platinum complexes. Nanoparticles and liposomes represent two major formulations that are in active clinical evaluation (32). The EPR effect allows significant increase in drug concentration; therefore, the cytotoxicity against tumor cells is increased, while normal tissues are spared from the drug-induced damage.

However, it is evident that the incomplete and immature vasculature within the tumors plays a fundamental role in drug resistance. The immature vasculature leads to reduced oxygenation and nourishment of cancer cells, and cancer cells adapt to grow in these critical conditions. The adaptation leads to changes in gene expression and metabolic pathways, which contributes to diminishing pH values in the tumor until an acidic pH is achieved and maintained (33). In these conditions, drug resistance phenomena may begin to occur because many drugs become ionized due to their pKa values within the range from 5.8 to 8.5 (34). Weak basic drugs, such as anthracyclines and vinca alkaloids, diffuse poorly in an acidic extracellular milieu because their ionized status obstructs their passage through cell membranes. Low pH may cause tumor resistance to mitoxantrone, a weak basic drug. Conversely, weak acid agents, such as chlorambucil and 5-fluorouracil, have an advantage in terms of distribution within the tumor and cytoplasmic sequestration because of the neutral-to-alkaline pH.
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Polymer micelles as powerful chemotherapeutic nanoparticles afford several advantages for targeted drug delivery in cancer, including increased drug solubility, prolonged circulation half-life, selective accumulation at tumor sites, and a decrease in toxicity. However, the technology still lacks tumor specificity and controlled release of the entrapped agents. Therefore, the focus has gradually shifted from passive targeting micelles to active targeting and responsive systems that carry additional mechanisms for site-specific release. pH-sensitive formulations are examples of how the versatility of micelles can lead to a fusion of chemical customization with biological insight to achieve active drug delivery.

In addition, hyperthermia may also increase drug accumulation within a tumor and has been evaluated in association with liposomes. Hypoxia may also exert a significant influence on drug sensitivity through the modulation of mRNA levels of several genes (35). For example, the chronic influence of hypoxia may lead to etoposide and vincristine resistance by modifying gene expression of HIF-1α (36). Furthermore, hypoxic conditions increase the heme content and induce the expression of ABCG2 protein in stem cells. This induction allows cellular survival by removing heme from the cytoplasm and thus diminishing the formation of reactive oxygen species (ROS) (37). Therefore, the hypoxia-induced expression of ABCG2 seems to give a double advantage to cancer cells, allowing survival in critical conditions and making the tumor resistant to drugs.

Modification of extracellular (pHe) and intracellular (pHi) could help to reverse drug resistance in tumors. Recent studies have demonstrated that some drugs exert their cytotoxic effects by altering the regulation of pHi through production of H2O2 in the mitochondria (38). Furthermore, an acidic pHi increases tumor sensitivity toward several drugs. Recent data support the use of proton-pump inhibitors (PPIs) to increase pHe and the pH of lysosomal organelles. Pretreatment with PPIs may reverse the MDR tumor phenotype, likely through the inhibition of drug excretion by ABC family members (i.e., ABCB1 or ABCG2). In some cases, it has been hypothesized that PPIs could induce drug accumulation within vesicle-like structures which cannot be excreted. Bioreductive drugs represent a logical consequence in the drug development process based on the knowledge of biological characteristics of tumors. In contrast with other bioreductive drugs, tirapazamine is active at intermediate oxygen concentrations, which acts synergistically with several antineoplastic agents such as cisplatin.

Block-copolymer micelles are spherical supramolecular assemblies of amphiphilic copolymers that have core–shell architecture. The core is a loading space that can accommodate hydrophobic drugs, and the shell is a hydrophilic brush-like corona that makes
the micelle water soluble. Block-copolymer micelles allow delivery of poorly soluble contents and avoid the pH e and pH i limitation. The micelles localize in several cytoplasmic organelles, including the mitochondria, but not the nucleus (39). Administering immunomicelles loaded with the anticancer drug taxol to mice with lung carcinoma resulted in increased accumulation of taxol in the tumor. Furthermore, nanoparticle shape may be important in designing better nanotechnology-based drug delivery vehicles. Filomicelles are about ten times longer than their spherical counterparts and are more persistent than any known synthetic nanoparticle (40). Preliminary results further demonstrate that filomicelles can effectively deliver the anticancer drug paclitaxel and shrink human-derived tumors in mice. Although these findings show that long-circulating vehicles need not be nanospheres, they lend insight into possible shape effects on nanoparticle function.

6. Nanoparticles Used for Tumor Treatment

Because cells will typically internalize nanomaterials below 100 nm, nanostructures have the ability to enter the cells due to their nanoscale size. Some of the leading nanostructures being used for this purpose include fullerenes, dendrimers, and nanoshells (Table 21.1). Fullerenes (or Buckyballs) are natural hollow spheres, 1 nm in diameter, made with carbon atoms. Fullerenes create a unique drug delivery platform that allows active pharmacophores to be conjugated to their surface in three-dimensional orientations for precise control in matching fullerene compounds to biological targets, in entrapping atoms within the fullerene cage, and for attaching fullerene derivatives to targeting agents. One of these fullerenes investigated by our group is \([\text{Gd@C}_{82}\text{(OH)}_{22}]_{n}\), which is a water-soluble hydroxyl modified metalfullerene. This nanoparticle has a strong capacity to enhance immunity and protect the normal tissues from tumor invasion, with almost no toxicity \(\text{in vivo}\) and/or \(\text{in vitro}\) (41). In comparison with conventional antitumor chemicals such as cisplatin and cyclophosphamide, this nanoparticle is highly efficient at suppressing tumor growth. Its action is not due to toxic effects on tumor cells because it does not affect tumor cell proliferation directly under the administrated concentration. The distribution in the tissues is mainly in bones (about 1% of administration), then the pancreas, kidney and spleen, in that order. About 50% of \([\text{Gd@C}_{82}\text{(OH)}_{22}]_{n}\) are excreted in the urine and 35% in the feces, which suggests that this nanoparticle reaches tissues and organs through blood circulation and does not remain in the blood after 24 h of administration. This could be improved by appropriate modification of this nanoparticle.
Another nanomaterial used for cancer treatment is the dendrimer, which was used to treat tumor cells without triggering an immune response. This is due to the dendrimer’s small size and branched structure. Dendrimers can be designed to release attached compounds in response to a specific molecule or chemical reaction. In addition, a layered sphere called nanoshell is being developed for cancer therapy. The nanoshell has a gold exterior layer which covers interior layers of silica and drugs. Nanoshells can be made to absorb light energy and then convert it to heat. As a result, when nanoshells accumulate next to a target area such as a tumor cell, they can release tumor-specific antibodies when infrared light is administered.

Successful design of nanoparticles to treat tumors effectively requires assembly of the appropriate targeting ligands on nanocarriers and long-circulating nanosystems with appropriate surface modification and the capability to control particle stability, aggregation, receptor binding and subsequent biochemical cascades and signaling processes. The size of the particles must be large enough (30–100 nm) to avoid leakage into blood capillaries but not so large (>100 nm) that they become susceptible to

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**Table 21.1**

<table>
<thead>
<tr>
<th>Subjects</th>
<th>Properties</th>
<th>Nanomaterials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nanocrystals</td>
<td>Materials with nanocrystalline structure are different in their atomic structure, crystallographic orientation, or chemical composition</td>
<td>Ceramic, metal (quantum dots, nanogold, nanosilver, etc.) and metal oxide nanoparticles (CuO, ZnO, TiO₂, SiO₂, etc.)</td>
</tr>
<tr>
<td>Carbon nanotubes/fullerenes</td>
<td>Carbon-based nanomaterials are composed mostly of carbon in the form of a hollow spheres, ellipsoids, or tubes etc. Fullerenes are characterized with an elongated sphere of carbon atoms formed by interconnecting six-member rings and 12 isolated five-member rings forming hexagonal and pentagonal faces</td>
<td>Hollow cylinders of carbon atoms such as: Carbon nanotubes (CNTs) Fullerenes including C60, C70, C80, Gd@C82, C84, etc.</td>
</tr>
<tr>
<td>Organic nanoparticles</td>
<td>Nanomaterials with three components: a central core, an interior dendritic structure (the branches), and an exterior surface (the end groups)</td>
<td>Polymers built from branched units with numerous chain ends on the surface such as various dendrimers</td>
</tr>
<tr>
<td>Inorganic–organic hybrid nanoparticles</td>
<td>Hybrid nanomaterials consist of one material as matrix filled with another material</td>
<td>Nanoparticles or nanofibers with at least two different materials such as polyhedral silicon-containing organic polymers</td>
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Other nanomaterials without clear definition, classification
macrophage-based clearance. Surface manipulation can control the particle aggregation at interstitial sites, optimizing nanoparticle retention at lymph nodes. Very small particles (1–20 nm) with long circulatory residence times could slowly penetrate the vasculature into the interstitial spaces, and be transported by lymphatic vessels to lymph nodes. This phenomenon is quite important when designing nanoparticles to allow differential leakage from the blood circulation system through the permeable endothelium in lymph nodes.

To date, many different nanoparticles have been synthesized and developed for effective treatment of tumors (Table 21.1). Recently, a multistage nanoparticle system has been employed. This multistage system consists of mesoporous material and nanoparticles as two major components. Modified mesoporous material is about to carry nanoparticles to their designated targeting site; it then degrades to release the nanoparticles into the targeted tissue. The released nanoparticles consequently merge into cells for efficient treatment.

7. Nanotechnology Can Improve the Bioavailability of Poorly Soluble Anticancer Drugs

Nanotechnology has been successfully utilized to create a new drug delivery system that can solve the problem of poor water solubility common to many promising and currently available anticancer drugs and thereby increase their effectiveness. The poorly soluble anticancer drugs require the addition of solvents in order for them to be easily absorbed into cancer cells. Unfortunately, these solvents not only dilute the potency of the drugs but increase toxicity as well. Silica-based nanoparticles are used to deliver hydrophobic anticancer drugs and other water-insoluble drugs to human cancer cells (42). The experimental results suggest that mesoporous silica nanoparticles might be used as a vehicle to overcome the insolubility problem of many anticancer drugs.

Paclitaxel is widely used to treat multiple types of solid tumors. The commercially available paclitaxel formulation uses cremophor/ethanol (C/E) as solubilizers. Other formulations including nanoparticles have also been introduced. The nanoparticle and C/E formulations showed significant differences when compared to paclitaxel itself. Tissue specificity of the two formulations was different too. The nanoparticles showed longer retention and higher accumulation in organs and tissues. The most striking difference was an eightfold greater drug accumulation and sustained retention in the kidney. These data indicate that the nanoparticulate formulation of paclitaxel affects its clearance as well as distribution in tissues with preferential accumulation in the liver, spleen, small intestine, and kidney.
As mentioned above, block-copolymer micelles with core-shell architecture provide a loading space that can accommodate hydrophobic drugs, and the shell is a hydrophilic brush-like corona that makes the micelle water soluble, thereby allowing delivery of the poorly soluble contents and accumulated in tumor. Other nanoparticles consisting of human serum albumin (HSA) and containing different antisense ODNs (ASOs) have also been used for drug delivery insoluble drugs (43). The preparation process was optimized regarding the amount of desolvating agent, stabilization conditions, as well as nanoparticle purification. Wartlick et al. found that the glutaraldehyde cross-linking procedure of the particle matrix was a crucial parameter for biodegradability and drug release of the nanoparticles (43). The drug loading efficiency increased with longer chain length and employment of a phosphorothioate backbone. It indicated that there was no cytotoxic effect observed under nanoparticle concentrations up to 5,000 µg/ml in different tumor cells. In this study, the entrapment of a fluorescent labeled oligonucleotide within the particle matrix was used to detect intracellular drug release of the carrier systems. It was revealed under confocal laser scanning microscopy that nanoparticles cross-linked with low amounts of glutaraldehyde could rapidly be degraded intracellularly and could lead to a significant accumulation of the ASO in cytosolic compartments of the tumor cells.

8. Resistance to Cisplatin: A Broadly Used Anticancer Drug

The platinum coordination complex known as Peyrone’s chloride was firstly synthesized and described by M. Peyrone in 1845; these findings were published in 1965 (44). In the 1960s, Barnett Rosenberg serendipitously discovered its chemotherapeutic cancer activity (45, 46). In 1968, following further tests against various bacteria, cisplatin was administered intraperitoneally to mice at the nonlethal dose of 8 mg/kg, and was shown to cause marked tumor regression (47). The patient was first treated with confirmatory in vivo tests performed by clinical testing in 1971. Cisplatin was approved by the US Food and Drug Administration (FDA) for clinical application in 1978.

Since the biological properties of cisplatin as a anticancer drug were accidentally discovered over 40 years ago, it has had a major impact on the chemotherapeutic treatment of various cancers and is still widely used today. Cisplatin is one of the most widely used and most effective cytotoxic agents, and is broadly employed in the treatment of epithelial malignancies such as lung, head and neck, ovarian, bladder and testicular cancer (48). The action mechanism of cisplatin involves covalent binding to purine DNA bases, which
primarily leads to cellular apoptosis. However, its continued clinical use is impeded by its severe adverse reactions including renal toxicity from renal tubular damage, gastrointestinal toxicity, peripheral neuropathy, asthenia, and ototoxicity (45, 46).

The major limitation in the clinical applications of cisplatin is the development of cisplatin resistance by tumors. This arises either by clonal expansion of tumor cells in the heterogeneous tumor cell population with inherent resistance to cisplatin (with mutations in specific genes that confer resistance), or by acquired resistance by some cells in the tumor during treatment and their clonal expansion after killing of the sensitive cells by the drug. Tumor proliferation could be mainly conferred by limited uptake of the cisplatin by drug-resistant cells. Much is currently understood about how tumors commonly exhibit resistance to cisplatin, either intrinsically or as acquired during the courses of therapy. Mechanisms explaining cisplatin resistance include the reduction in cisplatin accumulation inside cancer cells because of barriers across the cell membrane, the faster repair of cisplatin adducts, increased cytoplasmic detoxification and tolerance to DNA damage, the modulation of apoptotic pathways in various cells, the mislocalization of functional membrane protein and a higher concentration of glutathione and metallothioneins in some types of tumors (49). A number of experimental strategies to overcome cisplatin resistance are at the preclinical or clinical stages.

Reduced cisplatin intracellular accumulation is the common result in different types of cisplatin-resistant cell lines. Cisplatin is highly polar and enters cells relatively slow in comparison to other classes of small-molecule cancer drugs. The uptake of cisplatin is influenced by factors such as sodium and potassium ion concentrations, pH, and the presence of reducing agents. The role of transporters or gated channels has been postulated in addition to passive diffusion (50). So far, copper transporter-1 (CTR1) is considered to have a substantial role in cisplatin influx (51, 52). Loss of CTR1 was found to lead to a two- to threefold increase in drug resistance (53). In contrast to the mechanism of MDR, which is caused by the overexpression of ABC transporters, it is generally decreased uptake rather than increased efflux that predominates in cisplatin-resistant cells. The efflux proteins such as multidrug resistance protein-1 (MRP1, also known as ABCC1), MRP2 (also known as CMOAT or ABCC2) was reported to be partially associated with cisplatin resistance. The conjugation of cisplatin with glutathione was more readily exported from cells by the ATP-dependent glutathione S-conjugate export (GS-X) pump.
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...that is, MRP1 or MRP2 (54). Other studies also support a role for the glutathione metabolic pathway in acquired and inherited drug resistance to cisplatin.

Maintenance of cisplatin levels in tumors for prolonged periods is expected to eradicate cisplatin sensitive cells without offering them a chance to develop resistance. Hurdles are the side effects of cisplatin and the toxicity from the cumulative dose. Several mechanisms can contribute to cisplatin resistance. A common observation, repeatedly reported over many years in many tumor cells with acquired resistance to cisplatin, is that of reduced platinum accumulation in comparison to the parental cells (55). The reduction in cisplatin accumulation inside cancer cells because of the cell membrane barrier is currently considered a major mechanism of acquired cisplatin resistance (56). The copper transporter CTR1 appears to control the accumulation of cisplatin in Saccharomyces cerevisiae. CTR1-deficient cells have reduced the uptake of cisplatin, and are 1.9-fold more resistant to the cytotoxic effect of cisplatin (57). However, until recently, the underlying complex molecular mechanism by which cisplatin enters cells still remained poorly defined.

Drug delivery in cancer is important for optimizing the effect of drugs and reducing toxic side effects. Several nanobiotechnologies, mostly based on nanoparticles, have been used to facilitate drug delivery in cancer. The development of less toxic, nanoscale liposomal formulations of cisplatin has been hampered by the low water solubility and low lipophilicity of cisplatin, resulting in very low encapsulation efficiencies. Burger et al. reported a novel method to efficiently encapsulate cisplatin in a lipid formulation by repeated freezing and thawing (58). The method is unique in that it generates nanocapsules, which are small aggregates of cisplatin covered by a single lipid bilayer. The nanoparticles have an unprecedented drug-to-lipid ratio and an in vitro cytotoxicity higher when compared to free cisplatin. It suggests that the nanoscale encapsulation may also be generalized to other drugs with low water solubility and lipophilicity.

A polymer–metal complex formation between cisplatin and PEG-poly(glutamic acid) block copolymers were prepared before by Nishiyama et al. (59), and their utility was also investigated as a tumor-targeted drug delivery system. Cisplatin-incorporated micelles with a 28 nm size exhibited a sustained drug release and the decay of the carrier itself in physiological saline. These nanoscale micelles showed a remarkably prolonged blood circulation and effectively accumulated in solid tumor sites (59). These data suggest that micelles with cisplatin could be a promising formulation for the targeted therapy of solid tumors. Micelles with a hydrophobic inner core and hydrophilic outer shell allow the chemical entrapment of cisplatin into the micelles; cisplatin is then released slowly into the target organism. As the extracellular...
pH of solid tumors has often been shown to be more acidic than normal tissues, this might also explain, in part, the increased tumor delivery of micelles with cisplatin.

Overall, numerous mechanisms seem to be involved in tumor resistance to cisplatin studied in vitro. Studies have provided several rational approaches to circumventing clinical cisplatin resistance in patients. The strategy of using delivery vehicles to selectively transport more of a tumor-killing agent to tumors is attractive, and has now been clinically validated with the cytotoxics doxorubicin (liposomal doxorubicin) (60) and paclitaxel (nanoparticle albumin-bound paclitaxel) (61). To exploit the EPR effect of cisplatin in tumors, it has been linked to water-soluble, biocompatible nanomaterials. Trials are continuing with a reformulated cisplatin in an attempt to improve its antitumor activity.

Chemotherapy patients can be classified as either platinum-sensitive or platinum-resistant, depending on whether they have relapsed or progressed within 26 weeks of completing first-line platinum based chemotherapy (62). Expression of the mitogen-activated protein kinase phosphatase-1 (MKP-1) was a prognostic marker for patients with invasive ovarian carcinomas. The MKP-1 mRNA levels were strongly inducible upon treatment of OVCAR-3 cells with cisplatin. MKP-1 expression is a clinically useful marker to estimate patient prognosis as well as response to cisplatin chemotherapy.

Nanotechnology can be applied to encapsulate and protect drugs during transit in vivo. Drug encapsulation materials include liposomes and polymers (i.e., Polylactide (PLA) and Lactide-co-Glycolide (PLGA)). In addition to liposomes and polymers, other types of nanoparticles are also available for encapsulation. Materials such as silica and calcium phosphate (hydroxyapatite) have demonstrated superior properties at nanoscale rather than microscale, and can potentially be better suited for cisplatin delivery challenges. The materials form capsules around cisplatin and permit timed drug release to occur as the drug diffuses through the encapsulation material.

Lipoplatin is a liposomal cisplatin formulation currently under clinical trials. The advantage of lipoplatin appears to arise from its 2- to 50-fold higher concentration in human tumors when compared to normal human tissues in biopsies, measured as total platinum with atomic absorption. The lipoplatin formulation can attain a higher concentration in tumors via its preferential extravasation through altered and compromised tumor vasculature. In order to achieve this property, liposomes that enclose chemotherapy
drugs must have a diameter below 130 nm, long-circulation properties and the ability to escape immune surveillance.

The antitumor activity of cisplatin, encapsulated into transferrin-conjugated polyethylene glycol liposomes (Tf-PEG liposomes), was studied in nude mice with peritoneal dissemination of human gastric cancer cells. Small unilamellar Tf-PEG, PEG or DSPC/CH liposomes (bare liposomes) encapsulating cisplatin were prepared by reverse-phase evaporation followed by extrusion. The Tf-PEG liposomes were internalized into tumor cells by receptor-mediated endocytosis as shown by electron microscopy. Uptake of Tf-PEG liposomes into the liver and spleen was significantly lower than that of bare liposomes and had antitumor properties in nude mice xenografts that were better than free cisplatin (63).

A novel bile acid–cisplatin complex, called as Bamet-R2, with liver vectoriality, was synthesized with the aim of overcoming cisplatin resistance. This complex had increased water solubility by encapsulation into liposomes and enhanced uptake by liver tumor cells. Bamet-R2 was effectively incorporated into liposomes with an increase in the concentration of the drug by more than 6 million fold compared with that in the initial free solution; this is 1,000-fold higher than the encapsulation obtained for cisplatin (64). A lipophilic cisplatin derivative, NDDP, formulated in conventional liposomes was shown not to be a cross-resistant with cisplatin in different in vitro and in vivo systems, and more active than cisplatin against tumor metastasis (65). NDDP was also formulated in liposomes composed of phosphatidylcholine, cholesterol and monosialoganglioside or PEG conjugated to phosphatidylethanolamine with prolonged circulation (66).

Local and sustained release of cisplatin near or inside a tumor may have distinct advantages over systemic administration of the drug. Cisplatin formulations in gel-type materials suitable for intratumoral injection have been tested in several laboratories. In general, these methods suffer from inefficient loading of the drug and other hurdles relating to its release mode and overall toxicity of the formulation. Malignant bone tumors are treated with surgical therapy and simultaneous systemic chemotherapy. In order to overcome the toxicity of this approach, the bone-cementing apatite (calcium phosphate) was used for a cisplatin formulation to develop an implant and maintain high concentrations of cisplatin at local sites in animals to counteract local structural weakness after tumor resection and treat residual malignant bone tumors. Approximately 33% of the total bound cisplatin was released after
4.25 days. This approach might be used for the slow and local release of cisplatin in vivo (67, 68). PLGA-mPEG nanoparticles containing cisplatin were prepared by a double emulsion method and characterized with regard to their morphology, size, zeta potential and drug loading. Although intravenous administration of these cisplatin nanoparticles in mice resulted in prolonged cisplatin circulation in the blood, they suffered from loading efficiency for therapeutic applications (69). Degradable starch microspheres in an aqueous crystal suspension were used in clinical trials to achieve intensification of intraarterial chemotherapy of head and neck cancer with high-dose cisplatin (70).

Nanoparticle-formulated cisplatin might further broaden its applicability to tumor types such as prostate cancer and small-cell lung cancer. Improved tumor delivery strategies and controlled release of cisplatin with specific modulators of cisplatin-resistance mechanisms might also provide future clinical benefits. Such strategies are sometimes unsuitable for clinical practice because of technical and biologic constraints, but in some cases they represent fruitful efforts to improve cancer chemotherapy. In particular, novel pharmaceutical formulations of cisplatin improve treatment efficacy and tolerability by increasing drug delivery within tumors or in close proximity.

Nanotechnology provides a wide range of new technologies for developing customized solutions that optimize the delivery of pharmaceutical agents. To be therapeutically effective, drugs need to be protected during their transit to the target action site in vivo while maintaining their biological and chemical properties. Some drugs are highly toxic and can cause serious side effects and have reduced therapeutic effect if they decompose during their delivery. Once the drug arrives at its destination, it needs to be released at an appropriate rate so that it can be effective. If the drug is released too rapidly it may not be completely absorbed, or it may cause gastrointestinal irritation and other side effects. The use of nanoparticle for drug delivery could positively impact the rate of absorption, distribution, metabolism, and excretion of the drugs in the body. In addition, nanoparticle delivery can allow the drug to reach its target in a more active form. There are severe restrictions on the nanomaterials and synthesis processes that can be used in drug delivery systems. The drug delivery material must be compatible and has to be easily bound with the drugs; in addition, the nanomaterial has to be easily degraded after use. It can be either metabolized or eliminated via normal excretory routes.
Nanotechnology can offer new drug delivery solutions by drug encapsulation. When materials are encapsulated in nanoparticles within the 1–100 nm size range, they have a larger surface area for the same volume, smaller pore size, improved solubility, and different structural properties. This can improve both the diffusion and degradation characteristics of the encapsulated nanomaterial. Nanotechnology involves the creation and use of materials and devices at the atomic and molecular level. Because clinical chemotherapy uses a variety of molecular materials and devices, and nanotechnology has the potential to provide many medical and pharmaceutical insights, such as how molecular materials self-assemble, self-regulate, and self-destroy. The scope of nanotechnology is enormous and it overlaps with the traditional medicine. Although only a subset of nanotechnology is applied to biological processes including medical and pharmaceutical usage, the potential for breakthrough is enormous and is being pursued on multiple fronts.

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