TARGETING OF ANTI CANCER DRUGS THROUGH NANOPARTICLES

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ABSTRACT

Prior studies suggested that nanoparticle drug delivery might improve the therapeutic response to anticancer drugs and allow the simultaneous monitoring of drug uptake by tumours. Cancer Nano therapeutics are rapidly progressing and are being implemented to solve several limitations of conventional drug delivery systems such as nonspecific bio distribution and targeting, lack of water solubility, poor oral bioavailability, and low therapeutic indices. To improve the bio distribution of cancer drugs, nanoparticles have been designed for optimal size and surface characteristics to increase their circulation time in the bloodstream. They are also able to carry their loaded active drugs to cancer cells by selectively using the unique pathophysiology of tumours, such as their enhanced permeability and retention effect and the tumour microenvironment. In addition to this passive targeting mechanism, active targeting strategies using ligands or antibodies directed against selected tumour targets amplify the specificity of these therapeutic nanoparticles. Drug resistance, another obstacle that impedes the efficacy of both molecularly targeted and conventional chemotherapeutic agents, might also be overcome, or at least reduced, using nanoparticles. Nanoparticles have the ability to accumulate in cells without being recognized by P-glycoprotein, one of the main mediators of multidrug resistance, resulting in the increased intracellular concentration of drugs. Multifunctional and multiplex nanoparticles are now being actively investigated and are on the horizon as the next generation of nanoparticles, facilitating personalized and tailored cancer treatment. Targeting methotrexate increased its antitumor activity and markedly decreased its toxicity, allowing therapeutic responses not possible with a free drug.
KEYWORDS: Nanoparticles, Drug delivery, Targeting, Drug release, Anti-Cancer Drugs.

1. INTRODUCTION
Nanoparticles are defined as particulate dispersions or solid particles with a size in the range of 10-1000nm. The drug is dissolved, entrapped, encapsulated or attached to a nanoparticle matrix. Depending upon the method of preparation, nanoparticles, nanospheres or nanocapsules can be obtained. Nanocapsules are systems in which the drug is confined to a cavity surrounded by a unique polymer membrane, while nanospheres are matrix systems in which the drug is physically and uniformly dispersed. In recent years, biodegradable polymeric nanoparticles, particularly those coated with hydrophilic polymer such as polyethylene glycol (PEG) known as long-circulating particles, have been used as potential drug delivery devices because of their ability to circulate for a prolonged period time target a particular organ, as carriers of DNA in gene therapy, and their ability to deliver proteins, peptides and genes. The major goals in designing nanoparticles as a delivery system are to control particle size, surface properties and release of pharmacologically active agents in order to achieve the site-specific action of the drug at the therapeutically optimal rate and dose regimen. Though liposomes have been used as potential carriers with unique advantages including protecting drugs from degradation, targeting to site of action and reduction toxicity or side effects, their applications are limited due to inherent problems such as low encapsulation efficiency, rapid leakage of water-soluble drug in the presence of blood components and poor storage stability. On the other hand, polymeric nanoparticles offer some specific advantages over liposomes. For instance, they help to increase the stability of drugs/proteins and possess useful controlled release properties.

1.1. The advantages of using nanoparticles as a drug include the following
1. Particle size and surface characteristics of nanoparticles can be easily manipulated to achieve both passive and active drug targeting after parenteral administration.
2. They control and sustain release of the drug during the transportation and at the site of localization, altering organ distribution of the drug and subsequent clearance of the drug so as to achieve increase in drug therapeutic efficacy and reduction in side effects.
3. Site-specific targeting can be achieved by attaching targeting ligands to surface of particles or use of magnetic guidance.
4. The system can be used for various routes of administration including oral, nasal, parenteral, intra-ocular etc.
1.2. Cancer

Development of abnormal cells that divide uncontrollably which have the ability to infiltrate and destroy normal body tissue.

Cancer arises as a result of series of genetic and epigenetic changes

- Inactivation of tumor suppressor gene.
- The activation of oncogenes Cancer cells differs from normal cells.
- Uncontrolled proliferation.
- Ability to undergo metastasis.
- High interstitial pressure at the center.
- Solid tumours have a pore size of 100nm to 2um.

1.3. Preparation of Nanoparticles

Nanoparticles have been prepared most frequency by three methods:
(1) Dispersion of preformed polymers.
(2) Polymerization of monomers.
(3) Ionic gelation or coacervation of hydrophilic polymers.

However, other methods such as supercritical fluid technology and particle replication in non-wetting templates have also been described in the literature for production of nanoparticles. The latter was claimed to have absolute control of particle size, shape and composition, which could set an example for the future mass production of nanoparticles in industry (Van vlerken et al 2006).
1.3.1. Dispersion of preformed polymers
Dispersion of preformed polymers is a common technique used to prepare biodegradable Nanoparticles from poly lactic acid (PLA), poly glycolic acid (PLG), poly (D, L-lactide-co-glycolide) (PLGA) and poly cyanoacrylate (PCA), this technique can be used in various ways as described below (Tice et al 1985).

1.3.2. Solvent evaporation method
In this method, the polymer is dissolved in an organic solvent such as dichloromethane, chloroform or ethyl acetate which is also used as the solvent for dissolving the hydrophobic drug. The mixture of polymer and drug solution is then emulsified in an aqueous solution containing a surfactant or emulsifying agent to form an oil in water (o/w) emulsion. After the formation of stable emulsion, the organic solvent is evaporated either by reducing the pressure or by continuous stirring. Particle size was found to be influenced by the type and concentrations of stabilizer, homogenizer speed and polymer concentration. In order to produce small particle size, often a high-speed homogenization or ultrasonication may be employed.

1.3.3. Spontaneous emulsification or solvent diffusion method
This is a modified version of solvent evaporation method. In this method, the water miscible solvent along with a small amount of the water immiscible organic solvent is used as an oil phase. Due to the spontaneous diffusion of solvents an interfacial turbulence is created between the two phases leading to the formation of small particles. As the concentration of water miscible solvent increases, a decrease in the size of particle can be achieved. Both solvent evaporation and solvent diffusion methods can be used for hydrophobic or hydrophilic drugs. In the case of hydrophilic drug, a multiple w/o/w emulsion needs to be formed with the drug dissolved in the internal aqueous phase (Nah J.W.et al 2006).

1.3.4. Polymerization method
In this method, monomers are polymerized to form nanoparticles in an aqueous solution. Drug is incorporated either by being dissolved in the polymerization medium or by adsorption onto the nanoparticles after polymerization completed. The nanoparticle suspension is then purified to remove various stabilizers and surfactants employed for polymerization by ultracentrifugation and re-suspending the particles in an isotonic surfactant-free medium. This technique has been reported for making polybutyl
cyanoacrylate or poly alkyl cyanoacrylate nanoparticles. Nanocapsule formation and their particle size depends on the concentration of the surfactants and stabilizers used.

1.3.5. Coacervation or ionic gelation method
Much research has been focused on the preparation of nanoparticles using biodegradable Hydrophilic polymers such as chitosan, gelatine and sodium alginate. Calvo and co-workers developed a method for preparing hydrophilic chitosan nanoparticles by ionic gelation method. The method involves a mixture of two aqueous phases, of which one is the polymer chitosan, a di-block co-polymer ethylene oxide or propylene oxide (PEO-PPO) and the other is a polyanion sodium triply phosphate. In this method, positively charged amino group of chitosan interacts with negative charged triply phosphate to form coacervates with a size in the range of nanometre. Coacervates are formed as a result of electrostatic interaction between two aqueous phases, whereas, ionic gelation involves the material undergoing transition from liquid to gel due to ionic interaction conditions at room temperature.

2. TYPES OF TARGETING
2.1. ACTIVE TARGETING
Conjugating the nanoparticle to the targeted organ, tumor or individual cells for preferential accumulation. Active tumor targeting of nanoparticles involves attaching Molecules, known collectively as ligands, to the outsides of nanoparticles. These ligands are special in that they can recognize and bind to complementary molecules, or receptors, found on the surface of tumour cells.

Examples for ligands
- Folate, Biotin, Thiamine, Transferin (Feng et al 2003).
- Lecithin, Antibodies, Antibody fragments.
- Galactose, Apotamase.

2.2. PASSIVE TARGETING
The surface of the drug or polymer nanoparticle is coated with hydrophilic PEG or PLGA that reduces the hydrophobic interactions with the Reticulo Endothelial system (Krang D.M. et al 1995).
- The drug circulates in the blood for a long time.
- E.g. paclitaxel
Table-1: Different polymer-Drug conjugates using as Nanoparticles.

<table>
<thead>
<tr>
<th>S.No.</th>
<th>Polymer-drug conjugate</th>
<th>Disease</th>
<th>Company</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>PEG-Asparginases</td>
<td>Acute lymphopo blastic anemia</td>
<td>Enzon</td>
</tr>
<tr>
<td>2.</td>
<td>HPMA coplymer-Doxorubicin</td>
<td>Lung and breast cancer</td>
<td>Pfizer</td>
</tr>
<tr>
<td>3.</td>
<td>PEG-Paclitaxil</td>
<td>Clinical evaluation of solid tumours</td>
<td>Enzon</td>
</tr>
</tbody>
</table>

3. The nanoparticle drug delivery improve the therapeutic response to anti-cancer drugs.

3.1. Acetylated dendrimers were conjugated to folic acid as a targeting agent
Foliate receptor is over expressed in breast, ovary, endometrium, kidney, lung, head, neck, brain & myeloid cancers (Kommareddy et al 2005). E.g: methatrexate,

3.2. Twin nanoparticle shows effective at targeting, killing breast cancer cells
Treatment of breast cancer by twin nanoparticle by binding one gold nanoparticle with an iron oxide nanoparticle. A synthetic protein anti-body was attached to the iron oxide nanoparticle act as a targeting agent, on another end cis-platin was attached to the gold nanoparticle. The attached anti-body binds to the antigen protein located on the surface of the malignant cell, the drug release occurs at the malignant cell and kills it.

3.3. Carbon Nanotubes Target Tumor Cells
The Platinum-IV can incorporated into carbon nanotubes which has the capability to attach tumor targeting agent -Folic acid. The carbon nanotubes rapidly enter the target cell. There enzymes within the cell convert Platinum-IV to toxic Platinum-II which reacts with DNA and eventually kills the tumor cell (Hainfeld et al 2005).

3.4. Thermal cytotoxicity in malignant cells
The SWNTs targeted to cancer cells may allow noninvasive RF field treatments to produce lethal thermal injury to the malignant cells. The RF field can be used 13.56 megahertz.

3.5. Remote magnetic field triggers nanoparticle drug release
To the nanoparticle a short piece of DNA, one or more anticancer drugs were added. The nanoparticle and drugs are complimentarily attached to the DNA. At body temperature, the complimentary strands of DNA form the famous and strong double helix. When the
nanoparticle becomes warm as a result of an applied oscillating magnetic field the bonds of DNA become weak and drug molecule diffuses out.

3.6. Enhanced cytotoxicity of monoclonal anticancer antibody
Doxorubicin loaded long circulating liposomes were modified with the nucleosome specific monoclonal antibody 2c5 (mAb 2c5) as a result higher cytotoxicity towards various cancer cells (Esenaliev et al 2005).

3.7. Magnetic nanoparticle targeting human cancer cells
Nanoparticles are functionalized with ligands that bind with high affinity to the EphA2 receptor in the ovarian cancer.

4. APPLICATIONS OF NANOPARTICULATE DELIVERY SYSTEMS
4.1. The rationale of using nanoparticles for tumour targeting is based on
1. Nanoparticles will be able to deliver a concentrate dose of drug in the vicinity of the tumour targets via the enhanced permeability and retention effect or active targeting by ligands on the surface of nanoparticles.

2. Nanoparticles will reduce the drug exposure of health tissues by limiting drug distribution to target organ (Cho B.K. et al 1997).

3. Histological examination showed a considerable accumulation of nanoparticles in the liposomal vesicles of Kupffer cells, whereas nanoparticles could not be clearly identified in tumoral cells. Thus Kupffer cells, after a massive uptake of nanoparticles by phagocytosis, were able to induce the release of doxorubicin, leading to a gradient of drug concentration, favourable for a prolonged diffusion of the free and still active drug towards the neighbouring metastatic cells. When conventional nanoparticles are used as carriers in chemotherapy, some cytotoxicity against the Kupffer cells can be expected, which would result in deficiency of Kupffer cells and naturally lead to reduced liver uptake and decreased therapeutic effect with intervals of less than 2 weeks administration (Charlton D.E. et al 2006).

4. Moreover, conventional nanoparticles can also target bone marrow (MPS tissue), which is an important but unfavourable site of action for most anticancer drugs because chemotherapy with such carriers may increase myelosuppresive effect. Therefore, the ability of conventional nanoparticles to enhance anticancer drugs efficacy is limited to targeting tumours at the level of MPS-rich organs. Also, directing anticancer drug-loaded nanoparticles
to other tumoral sites is not feasible if a rapid clearance of nanoparticles occurs shortly after intravenous administration.

4.2. Long circulating nanoparticles
To be successful as a drug delivery system, nanoparticles must be able to target tumours which are localized outside MPS-rich organs. In the past decade, a great deal of work has been devoted to developing so-called “stealth Nanoparticles”.

- Particles or PEGylated nanoparticles, which are invisible to macrophages or phagocytes major breakthrough in the field came when the use of hydrophilic polymers (such as polyethylene glycol, poloxamines, poloxamers, and polysaccharides) to efficiently coat conventional nanoparticle surface produced an opposing effect to the uptake by the MPS.
- These coatings provide a dynamic “cloud” of hydrophilic and neutral chains at the particle surface which repel plasma proteins.
- As a result, those coated nanoparticles become invisible to MPS, therefore, remained in the circulation for a longer period of time. Hydrophilic polymers can be introduced at the surface in two ways, either by adsorption of surfactants or by use of block or branched copolymers for production of nanoparticles.
- Studies show nanoparticles containing a coat of PEG not only have a prolonged half-life in the blood compartment but also be able to selectively extravasate in pathological sites such as tumours or inflamed regions with a leaky vasculature.
- As a result, such long-circulating nanoparticles have increased the potential to directly target tumors located outside MPS-rich regions.
- The size of the colloidal carriers as well as their surface characteristics are the critical to the biological fate of nanoparticles. A size less than 100 nm and a hydrophilic surface are essential in achieving the reduction of opsonisation reactions and subsequent clearance by macrophages.
- Coating conventional nanoparticles with surfactants or PEG to obtain a long-circulating carrier has now been used as a standard strategy for drug targeting in vivo.
- Extensive efforts have been devoted to achieving “active targeting” of nanoparticles in order to deliver drugs to the right targets, based on molecular recognition processes such as ligand-receptor or antigen-antibody interaction. Considering that fact that folate receptors are over expressed on the surface of some human malignant cells and the cell adhesion molecules such as selectins and integrins are involved in metastatic events, nanoparticles bearing specific ligands such as folate may be used to target the cells.
• Ovarian carcinoma while specific peptides or carbohydrates may be used to target integrins and selectins (Brannon et al 2004).

4.3. Nanoparticles for oral delivery of peptides and proteins

• Significant advances in biotechnology and biochemistry have led to the discovery of a large number of bioactive molecules and vaccines based on peptides and proteins. Development of suitable carriers remains a challenge due to the fact that bioavailability of these molecules is limited by the epithelial barriers of the gastrointestinal tract and their susceptibility to gastrointestinal degradation by digestive enzymes. Polymeric nanoparticles allow encapsulation of bioactive molecules and protect them against enzymatic and hydrolytic degradation. For instance, it has been found that insulin-loaded nanoparticles have preserved insulin activity and produced blood glucose reduction in diabetic rats for up to 14 days following the oral administration.

4.4. Targeting of nanoparticles to epithelial cells

In the GI tract using ligands

• Targeting strategies to improve the interaction of nanoparticles with absorptive enterocytes and M-cells of Peyer’s patches in the GI tract can be classified into those utilizing specific binding to ligands or receptors and those based on nonspecific mechanism (Wiener EC et al 1997).

• The surface of enterocytes and M cells display cell-specific carbohydrates,

Which may serve as binding sites to colloidal drug carriers containing appropriate ligands.

Certain glycoproteins and lectins binds selectively to this type of surface structure by specific receptor-mediated mechanism. Different lectins, such as bean lectin and tomato lectin, have been studied to enhance oral peptide adsorption.

• Vitamin B-12 absorption from the gut under physiological conditions occurs via Receptor-mediated endocytosis. The ability to increase oral bioavailability of various peptides (e.g., granulocyte colony stimulating factor, erythropoietin) and particles by covalent coupling to vitamin B-12. For this intrinsic process, mucoprotein is required, which is prepared by the mucus membrane in the stomach and binds specifically to
cobalamin. The mucoprotein completely reaches the ileum where resorption is mediated by specific receptors.

4.5. Nanoparticles for drug delivery into the brain

The blood-brain barrier (BBB) is the most important factor limiting the development of new Drugs for the central nervous system. The BBB is characterized by relatively impermeable Endothelial cells with tight junctions, enzymatic activity and active efflux transport systems. It effectively prevents the passage of water-soluble molecules from the blood circulation into the CNS, and can also reduce the brain concentration of lipid-soluble molecules by the function of enzymes or efflux pumps consequently, the BBB only permits selective Transport of molecules that are essential for brain function (Bhadra et al 2002).

5. CONCLUSION

The foregoing show that Nano particulate systems have great potentials, being able to convert poorly soluble, poorly absorbed and labile biologically active substance into promising deliverable drugs. The core of this system can enclose a variety of drugs, enzymes, genes and is characterized by a long circulation time due to the hydrophilic shell which prevents recognition by the reticular-endothelial system. To optimize this drug delivery system, greater understanding of the different mechanisms of biological interactions, and particle engineering, is still required. Further advances are needed in order to turn the concept of nanoparticle technology into a realistic practical application as the next generation of drug delivery system.

6. REFERENCES