DRUG TARGETING IN NEOPLASTIC DISORDERS

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ABSTRACT

The review on drug targeting in neoplastic disorders emphasizes the various targeting techniques for effective treatment of the condition. Neoplasm refers to the new growth and its common in any individual, uncontrolled neoplasm results in condition called neoplastic disorder or tumor or cancer. Globally 14.1 million cancer cases were reported in 2012 and the number is estimated to reach 24 million by 2035. This review outlines the advances in drug targeting to the neoplastic disorders keeping in view on the problems associated with conventional chemotherapy and radiation therapy. It highlights the novel drug delivery systems like liposomes and particulate system like nanoparticles usage in treatment of cancer and effective targeting ways to target the tumor cells without affecting the normal cells.

Keywords: Chemotherapy, Liposomes, Nanoparticles, Neoplasm.

INTRODUCTION

As the evolution of science is advancing the advancement of diseases are also progressing. Worldwide cancer figures as one of the major cause of the morbidity and mortality according to World Health Organization1. Globally 14.1 million cancer cases were diagnosed in 2012 and the number is estimated to reach 24 million by 20352. Around 8 million deaths were reported in 20121. Neoplasm refers to the new growth of cells which is a common mechanism involved in the anabolic processes and for the survival of living organisms. Growth among the cells is primarily regulated by two genes i.e. oncogenes and tumor suppressor genes. Any loss in the regulatory mechanism of these genes will result in uncontrolled neoplasm which is commonly referred as neoplastic disorder or cancer or tumor. Cancer may be formed in any part of the body and it is mostly observed in lungs, breast and prostate according to World Cancer Research Fund International2.

CELLULAR GROWTH-CANCER FORMATION

Cellular growth is an intrinsic mechanism for multiplication of cells and it is majorly regulated by two important classes of genes3. They are oncogenes and tumor suppressor genes. Oncogenes or proto-oncogenes also help in the promotion of cellular growth by coding of growth stimulatory protein whereas tumor suppressor genes are useful to inhibit the growth of the cells. Prior to cell proliferation the cell enlarges and divides and for the survival of a cell it is needed to get attached among them or to the extra cellular matrix, cell adheres to the extracellular matrix with the aid of area codes. If cell adhesion does not takes place cell may undergo apoptosis which results in death of cell. Any of the mutation in either of the genes results in excessive multiplication of the cells which is a neoplastic disorder or commonly called as cancer. Cancer can also be caused due to replication errors, exposure to chemical carcinogens, radiations and sometimes also due to viral infections.

TYPES OF CANCERS

Cancers are of two types, they are benign tumor and malignant tumor. Among these benign tumor does not migrate to other place from the place of formation and they can be removed by
surgery, where as malignant tumors are having capacity to migrate to other parts of body which is primary danger with malignant tumors. If the illness reaches to any of the body’s vital organs it would be fatal sometimes. Malignant tumors cannot be removed easily by surgery and reoccurrence is seen even on removal of these types\(^5\). They need to be treated by chemotherapy, radiation or by other means.

**SUB-TYPES OF CANCER**

1. **Carcinoma**: Cancers arising from epithelial tissues like breast, body cavity, skin and prostate.
2. **Sarcoma**: Cancers arising from supporting tissues or any connective tissues like blood vessels, bone, cartilage, fat, muscles and nerves.
3. **Lymphomas & Leukemias**: Cancers arising from blood-cells forming tissues i.e. lymph nodes, spleen & bone marrow.

**STEPS INVOLVED IN CANCER FORMATION**

Initially tumor cell evolution takes place due to any of the genetic mutation, the genetically altered cell and its off springs seem to be normal, but reproduce excessively which is referred as hyperplasia. The offspring of cell proliferates and appear abnormal in shape called as dysplasia. The genetically altered cell continues to grow abnormally and if tumor mass does not invade to other parts, it is called benign tumor or in-situ tumor. When the tumor mass invades underlying tissue and moves into blood or lymph and generate blood supply for the newly formed cells it may migrate to other parts of the body called as malignancy. The tumor mass develops blood supply by the process called angiogenesis. Cancer cells differ from normal cell in adhesion; they even survive for years if not adhere among themselves or to the extracellular matrix without undergoing cell suicidal or apoptosis. This is a challenging aspect in targeting of drugs to tumor.

**BARRIERS OFFERED BY TUMOR**

Lack of tumor cellular adhesion will result in no recruitment of leukocyte cells at the inflamed area during inflammation; by this tumor cells are avoiding attack of immune cells which is a part of host’s defense mechanism. The basement membrane of tumor vasculature varies. It is continuous in some areas, fenestrated with fenestrations of 30-80 nm diameters in places like liver, spleen and in bone marrow, whereas it is discontinuous with large fenestration of 100 nm of more in some parts of lungs, liver and spleen. In tumor vasculature basal lamina or basement membrane is helpful for lining and support of endothelium, so that in cancer chemotherapy the drug or drug carrier have to cross endothelial linings of blood vessel, from there have to reach interstitial fluid, then to extra cellular matrix and finally to tumor cells. The barriers like basement membrane, endothelial linings, and angiogenesis related factors; high interstitial pressure in the tumor vasculature, presence of p-glycoprotein pump clearance contributes to poor delivery of drug and targeting to the tumors.

**CONVENTIONAL THERAPY**

The conventional therapeutic methods include chemotherapy and radiation therapies. Chemotherapy includes administration of anti-neoplastic agents which kills tumor cells and also all actively growing cells which results in gastric ulceration and bleeding, anemia, alopecia, destruction of regenerated cells of lungs, liver and maintenance of higher concentration of drugs in plasma which results in unwanted affects and adverse effects of the drugs. Radiation is given directly to the site of tumor which easily damages the DNA of cells, exposure to the radiation may also affects some of the non-cancerous tissues.

**NOVEL DRUG DELIVERY SYSTEMS**

These are tailor made systems to overcome disadvantages associated with the conventional therapeutic systems. These systems also aim to identify the possible novel strategies for efficient and effective targeting of drugs in lower concentrations over a longer period of time within the therapeutic range directly to the cancer cells without affecting the normal cells\(^5\).

**MOLECULAR TARGETS FOR TUMOR THERAPY\(^3\)**

During cancer development various receptors, ligands, epitopes and surface determinants are over expressed on the tumor vasculature. Recognition of these will help to achieve targeting without affecting the normal cells. These are developed majorly in the tumor vasculature due to tumor progression and are absent in the normal cell growth. Some of them are expressed during certain stages of cellular differentiation; examples include transferrin receptor (Tfr), folate receptors, epidermal growth factors; and various immunological determinants.
factor receptor (EGFR), apo-lipoprotein receptor and haemopxin receptor. Surface determinants like Ia antigens, tumor associated antigens (Taa) are expressed exclusively on malignant cells. Epitopes in the form of angiogenic peptides which are formed during the angiogenesis is helpful to target the vascular sprouts of blood vessels and proliferating tissues.

STRATEGIES FOR TUMOR TARGETING

Site specific drug delivery presumes localization of drug and its carrier within the target organ, recognition and simultaneous interaction of carrier with specific target cells followed by delivery of therapeutic concentration of drug to the target cells with a little or no uptake by the non-target cells.

Targeting is of two types: 1. Active targeting & 2. Passive targeting

Active targeting is done by altering natural distribution pattern of drug carrier whereas passive targeting can be done through natural distribution pattern of drug carrier.

SITE SPECIFIC DRUG DELIVERY

Site specific drug delivery is classified into three types according to the specificity levels achieved in the delivery process:

I. Organ Targeting: drugs are delivered to the individual organs or tissues.

II. Cellular Targeting: drugs are delivered to the specific cell type within a tissue.

III. Intracellular Targeting: drugs are delivered to the different intracellular compartments in the target cells by special transport pathways.

The strategies of tumor targeting are discussed as follows:

1. Immunotherapy strategies
2. Multi-drug resistance targeting
3. Novel drug delivery systems
4. Other targeting strategies

1. IMMUNOTHERAPY STRATEGIES

Immunotherapy comes under passive targeting; it is aimed to attack disease with defense mechanism of body by using tumor antigens, antibodies, recombinant antibodies and monoclonal antibodies.

a. Tumor antigens: These are popularly known as tumor vaccines, they exhibit prompt immune response in individuals by activating their own immune system. Whole tumor cell can be used for this purpose. It can be given in extract form or whole cell can be used in an inactivated form. Nucleic acids like DNA & RNA coding for tumor antigens can also be used. Vaccines are having the advantage of showing reproducible production of antibodies against the tumor forming cells on subsequent attacks and also boost up the immune system.

b. Antibodies: They bind to the antigens of tumor cells and by this they are susceptible to destruction by the host’s immune system. They are further helpful in targeting blood vessels that supply to tumor, also to target its connective tissues. They can be used as a guided missile to recognize and deliver the therapeutic compounds to the tumor sites. Antibodies also acts as toxins which inhibit protein synthesis in tumor cells and in turn inhibit the tumor growth, they also trigger cytokines & inflammatory mediators which are helpful in these cell destruction.

c. Antibodies with prodrugs: When prodrugs are attached to tumor specific antibodies, they are converted to active moiety by enzymatic action and they show action against the tumor cells. Antibodies often linked directly with chemotherapeutic agents by which direct targeting of agent to the tumor cell can be achieved and it reduces associated side effects and toxic effects. Advancement in usage of antibody is antibody directed enzyme prodrug therapy (ADEPT), it utilizes the principle of localization of intravenously administered antibody with enzyme as a conjugate in the tumor tissue and the enzyme activates the prodrug within the tumor.

d. Recombinant antibodies: Recombinant antibody fragments can be used for development of anti-tumor strategy. Generally immunoglobulin on the surface of target cell exposes its tail region (Fc) for recognition by Fc receptors phagocytic cells of mono nuclear phagocytic system helps in tumor cell killing with the help of antibodies developed against the tumor surface antigen. Instead of using complete immunoglobulin, a fragment with antigenic binding property can be used so that the molecular size decreases by two times.
e. **Monoclonal antibodies**: Monoclonal antibodies also show promising delivery of drugs to the target cells, they are the antibodies obtained by hybridoma technology which are having the affinity to bind specifically to the desired antigen. By coupling of anti neoplastic agents to the monoclonal antibodies, the drug is delivered to the target site. The problems associated with monoclonal antibodies are,

- Slow elimination of these from blood and heterogeneous tumor uptake
- Poor vascular permeability
- Cross reactivity of normal tissues with monoclonal antibodies
- Metabolism of monoclonal antibodies and its conjugates
- Immunogenicity with murine forms in humans

Other strategies in immunotherapy are by immunotoxins, cytokines like Interleukin-2 and they enhance the ability of T-cells of immune system to recognize tumor cells. Tumor peptides, fragments of tumor proteins can also be utilized for achieving the targeting to immune system; they can be administered alone or with an adjuvant.

2. **MULTI DRUG RESISTANCE TARGETING**

Multi drug resistance is the problem associated with chemotherapy, the initial and subsequent chemotherapy allows tumor to develop secondary resistance. After getting resistance by the tumor cells additional chemotherapy will induces toxic effects in subjects without reducing the tumor growth. Tumor cells with multi drug resistance over expresses an energy dependent drug transport protein commonly referred as p-glyco protein (p-gp). This over expression leads to lowering of drug accumulation within the tumor cells since the cell is having ability to pump out hydrophobic anti tumor drug molecules with p-glyco protein pump. P-gp further detects and expels the drug and it just acts like a hydrophobic vacuum cleaner. Specific class of drugs known as multi drug resistance reversal agents can be used to overcome these problems, eg: Doxorubicin. These reversal agents acts by binding to P-glycoprotein and they block this channel, so that the drug is allowed to slip into the tumor cell and they can inhibit or kill the tumor cells.

Multi drug resistance reversal agents are often used in conjunction with liposomes for appropriate targeting due to the below stated reasons.

- Negatively charged phospholipids like phospholipids or cardiolipin in liposomes directly regulate the p-gp transporter.
- Liposomes provide sustained high levels of drug to the resistant cells over longer periods of time.
- After endocytosis of drug loaded liposomes by lysosomes, lysosomes protect liposomes from action of p-gp and avoid immediate contact with the transporter located at the plasma membrane.

3. **NOVEL DRUG DELIVERY SYSTEMS**

They are the promising drug delivery system that provide site specific targeting of drug to the tumor cells, elicits therapeutic concentration of drug at the effected region without affecting the normal system. They include vesicles like liposomes, niosomes, particulate systems like microparticles and nano particles. The primary aim of these systems is to ensure safety and to improve efficacy along with patient compliance. Of all novel drug delivering systems liposomes are widely selected for targeting in neoplastic disorders owing to its smaller size, compatibility with tissues and many more advantages.

a. **LIPOSOMES**

They are helpful for passive targeting to different tumors because of their longer circulatory half lives and they moves into tissues easily due to their vascular permeability due to low size and lipid nature. Specially engineered liposomes called stealth liposomes or pegylated liposomes which are stearically stabilized and coated by polyethylene glycol are used for longer circulation and increased permeability for movement into the vascular endothelium of the tumor cells.

On the other hand liposomes can be used in active as well as in passive targeting to the affected cells. Passive targeting can be achieved by using conventional liposomes and immunoliposomes. Immunoliposomes are produced by attaching liposomal surface to the antibodies developed against the tumor cells or against the specific surface antigens present on the tumor vascular endothelium. Active targeting can be achieved by usage of ligand mediated targeting in which ligands are developed against cell receptors or antigenic determinants which are expressed exclusively on the tumor vasculature. The ligands include antibodies,
glycolipids, glycol proteins, polysaccharides, proteins and immune regulatory molecules. Physicochemical strategies can also be utilized for targeting of liposomes, in this drug-carrier is given to the body and they release the drug only when exposed to specific micro environments such as change in pH, temperature or subjecting to the external condition like light, magnetic fields. Examples of liposomes targeting by physicochemical strategies include pH sensitive liposomes, thermo sensitive liposomes, photo activated liposomes and magneto liposomes.

**pH sensitive liposomes**
The pathological tissues of tumors have ambient pH which is considerably lower than the normal tissue. pH sensitive immuno liposomes can deliver the drugs directly to the cytoplasm of the tumor cells. Eg: dyes like calcininin, drugs like methotrexate.

**Thermo sensitive liposomes**
Thermo sensitive liposomes are positively charged liposomes having encapsulated paramagnetic materials often referred to as magneto-cationic liposomes. They release the drug on application of external heat, these thermo sensitive liposomes used as effective tools for hyper thermic treatment of solid tumors.

**Photo activated liposomes**
Photo activated liposomes consist of photo sensitive polymer in the external surface of liposomes and they release the drug from the liposomes on external application of light and they are helpful for the localized delivery of drug to the tumors.

**Magneto liposomes**
Magneto liposomes are having iron containing particles like magnetite in the liposomal surface. They release the drug on external application of magnetic field. This strategy is helpful in localization and controlled delivery of drug to the tumors. Magneto responsive thermo sensitive liposomes are modification of magneto liposomes and they release the drug in response to magnetic modulation and hyper thermia. Conventional liposomes are up taken by Reticulo Endothelial System (RES) and they are used to target the reticulo endothelial system organs like liver, lungs and spleen, whereas stealth liposomes are used to target non-reticulo endothelial system organs and they are having longer circulatory half-life since they are having capability to escape the circulating scavengers and degradation by these organs.

**b. NIOSOMES**
Niosomes are the vesicles similar to liposomes and along with lipid a non-ionic surfactant is employed in this formulation. They are having similar properties of entrapment of drugs and they are stable compared to other formulations and they show varied drug distribution characters and release characters since the surfactant used is of non biological origin and it escapes clearance of phagocytic system and shows a prolonged release characters.

**c. PARTICULATE SYSTEM IN CHEMOTHERAPY**
It is the most promising application and by this we can achieve targeting to the tumor tissues. Particulate system includes nanoparticles and microspheres. By intravenous administration accumulation of these particulate systems in the tumors is favored. The reason beyond this is presence of enhanced endocytic activity of immune system and leaky vasculature of the tumor cells. Stealth microparticles or nanoparticles can be prepared by coating them with polyoxyethylene or with dialkyl polyoxyethylene and phospholipids they are used to target the non reticulo endothelial system organs whereas non stealth or conventional microparticles or nanoparticles can be used to target the reticulo endothelial systems. Polymers like poly lactide, poly glycolide, poly lactide-co glycolide and polyvinyl pyrrolidone (PVP) are used in nanoparticles preparation along with polyalkyl cyano acrylates, eg: Taxol in PVP nanoparticles. Particulate system can be used for site specific targeting by coating them with antibodies. In this anchoring of target specific antibodies to the particulate system is more helpful in targeting. Monoclonal antibodies can be fixed on to the surface of particulate system by direct absorption or by covalent linkage. Following are the examples of drugs that are targeted by nanoparticles.

- Doxorubicin in polyisohexyl cyanoacrylate nanoparticles
- Doxorubicin in polyalkyl cyanoacrylate nanoparticles
- Mitoxantrone in polybutyl cyanoacrylate nanoparticles
4. OTHER TARGETING STRATEGIES
Other targeting strategies include targeting the angiogenesis, usage of chemo embolization and by receptor level targeting.

ANGIOGENESIS
Angiogenesis means formation of new capillaries by the tumor cells for their existence, as the tumor mass increases the new capillaries starts in the form of vascular sprouts and gradually develops into new capillaries. This process is also called as neovascularization. It continues between cells and passes to the other parts also, the danger associated with angiogenesis is that malignancies pass over the body by these newly formed capillaries and may enter the systemic circulation which might migrate to the vital organs of the body. Angiogenesis inhibitors like thrombospodin & angiostatin can be used to overcome the angiogenesis. They inhibit the new blood vessel formation, limits the tissue perfusion, inhibits vascular sprout generation, also interfere with endothelial growth factors which stimulates tumor growth.

CHEMO EMBOLIZATION: INTRA-ARTERIAL INFUSION CHEMOTHERAPY
In this chemotherapeutic agent with novel drug carriers like microcapsules and liposomes can be used. Implantable equipment can be used for intra arterial use. In this biodegradable particles are administered to target to the liver tumors i.e. drugs are given in an injection form with the help of a catheter into the artery that directly leads to the tumor. This can be used for continuous infusion or for a single use. The solution on reaching the target as a bolus localizes in the target organ and shows reduction in the tumor growth. This way is helpful to treat the patients for a longer time and if needed more frequently with less discomfort.

RECEPTOR LEVEL TARGETING
Receptor level targeting can be attained by targeting through the altered or over expressed receptors i.e. through folate, transferrin, lectin & lipoprotein receptors.

Targeting through folate receptor
Folate receptor is a highly selective marker over expressed in many of the ovarian carcinomas and epithelial tumors. These receptors can be targeted by using monoclonal antibody linked to folic acid, which is a high affinity ligand. Low molecular weight radio pharmaceuticals conjugated with folate ligand is tested in animal tumor model which is under study. The smaller size, wider availability, ease of conjugation, lack of presumed immunogenic property confirms folic acid as an ideal ligand in targeting.

Targeting through transferrin receptor
Transferrin receptor is also over expressed on some of the tumor cells and this can be utilized for targeting. Attachment of transferrin in delivery of drugs like methotrexate showed superior binding and subsequent reduction in tumor size as compared to the conventional carrier systems. Receptor targeted liposomes with transferrin is under investigation for gene therapy of tumor cells.

Targeting through lectin receptor
Lectin receptor is showing up regulation as well as down regulation in normal cells and also in tumor cells of liver. The reason is that they might be depleted in normal cells as well as in tumor cells due to the cellular redistribution, hence lower level of targeting was anticipated in patients with malignancies related to liver.

Targeting through lipoprotein receptor
Tumor cells also express receptor for low density lipoprotein which can be used as a targeting strategy. The number of these receptors in normal cells is regulated by metabolic factors like intracellular amount of cholesterol. In case of tumors these cells over express lipoprotein receptors whereas its number in normal cells declines. So it can be used as an effective targeting tool. For this targeting anti neoplastic drugs as well as photosensitive porphyrins can also be used. The porphyrin along with drug is loaded into a lipid vesicle like liposome, administered into localized tumor and subsequent exposure of irradiation or visible light resulted in the release of drug at the target site, since light exposure causes changes in porphyrin i.e. conversion of light energy to chemical energy and it resulted in severe biological damage of the target tissue. The disadvantage associated in this targeting is that solid tumors are having less access to lipoproteins.

FUTURE PERSPECTIVES
Vascular targeting can also be done with the help of phage peptides; they are capable of attaching to the developing tumor tissues. These peptides bind to the receptors that are present in the tumor angiogenic neovasculature & vascular...
targeting is helpful for the inhibition of angiogenesis and in turn helps for site specific targeting; eg. Doxorubicin targeting to tumor angiogenic vasculature by use of phage peptides. Peptide nucleic acids (PNA) is a DNA mimic with pseudopeptide backbone composed of aminoethyl glycine units, research scientists expressed usage of PNA’s will be helpful in the gene targeting of tumor cells.

CONCLUSION
The current review highlights various modalities which are under study to overcome the problems associated with conventional therapies, focuses on selective targeting of drug moiety to the desired affected tumor cells. A more thorough in-depth study regarding uptake by various tumor cells and their expressions is needed for refinement of the targeting strategies. Targeting by novel carriers deserves the attention as such carriers offers the possibility of targeting a wider range of therapeutic systems. Further a combination therapeutic regimen utilizing advantages of localized or direct administration and tumor specific delivery systems is of significant importance to the cancer patients. Recombinant technologies with the bio-pharmaceutical industry is producing various vaccines and is studying gene level targeting in tumor therapy which will be a trend setting in the modern pharmaceutical era.

REFERENCES