SMART Drug Based Targeted Delivery: A New Paradigm for Nanomedicine Strategies

Abstract

Introduction: Targeted drug delivery systems are nanoscale drug carrier molecules designed for improving the communication of cellular and molecular components and biodistribution of tumour targeted drug (chemo) therapeutics. Nanomaterials are generally clusters of molecules, atoms and molecular fragments into extremely small size particles (1-100 nm) in nature. Nanomaterials engineered as self-assembled biodegradable particles were used for targeted drug delivery system. Nanocarriers/particles should be capable of transporting high doses of chemotherapeutic drugs/nanomedicines into the targeted tumor cells without disturbing the normal healthy cells. It is also used for construction of novel targeted drug delivery system and future application in nanovaccination and nanotechnology.

Conclusion: Multifunctional smart nanoparticles or carries hold out the possibility of effective drug targeted therapeutics in molecular and cellular levels at the earliest stage. Here, we briefly discuss the significance of targeting strategies and drug delivery system and outline the current approaches and future directions in the improvement of tumor targeting nanomedicines.

Discussion

Ideal design for smart drug (Nanomedicine)

Smart drug which are ideally designed to satisfy the requirements will pave the way for targeted drug delivery. These requirements includes: a) high stability (longevity) and not easily degraded or cleared by the reticuloendothelial system (RES) during blood circulation, b) accumulation of high concentration of drug at the tumor targeted sites or area, c) effective intracellular drug delivery at the tumour targeted area or site that matches drug pharmacodynamics of kinetics and spatial control and d) tolerability. The blood circulation kinetics and spatial control and d) tolerability. The blood circulation and chemical nature of the encapsulated drug and encapsulation techniques. The design of tumor drug targeted delivery system based on: passive drug targeted delivery as a significance of enhanced permeability and retention effect (EPR) (without ligand targets and active drug release mechanisms) [11,12], targeted drug release based on recognition of receptor molecules [13-16], triggered drug release or stimuli – responsive release (pH, redox potential, temperature, light intensity, ROS response magnetic field, hypoxia response, glucose

Nanotechnology has led to a connection of various fields including applied physics, chemistry, optics, biology, computational modelling and analysis, pharmaceuticals and material science. Nanoscales construction provides the ability to analyse and manipulate the stimulation, target the behaviour of biological system or predict the structure by molecular modelling and computational science [1]. Side effects should be avoided by early diagnosis and effective treatment by using nanomedicines [2]. Nanotechnology in health-care research can positively result in enormous health benefits [3] collectively increasing research funding from all over the world [1]. Impacts from improvements progressing from nanotechnology developments over the next 20 years have been assessed to be nearly $1 trillion by studies carried out at the International Science Foundation [3]. Nanodevices or sensors and nanomaterials which are used for reparation, monitoring, construction and control of human biological system at cellular and molecular level were known as nanomedicines [3]. Administration of drug by transdermal, oral, intravenous and trans epithelial delivery system is fulfilled using nanocarrier as vehicles. Nanocarriers are present in different forms including polymeric, metal, liposomes, carbon, magnetic and biological particles [4–10] which have a huge biomedical application on drug delivery systems.

Although few products are awaiting for clinical trials, but still nanomedicines are found to be specially promising application and at present used in various fields of diagnosis of diseases, molecular imaging, delivery of chemotherapeutic drug to its targeted tumor sites. Nanomedicines play a significant role by enhancing the efficacy of therapeutic index and safety measurements of new targeted therapeutics even at low dosage levels [9]. Nanocarriers structures which holds conjugated drugs, encapsulated, adsorbed, dispersed and imaging agents. Nanovehicles including nanoemulsions liposomes, micelles, nanoparticulate system (polymer, carbon, drug nanoparticles, ceramic-based, biological, metal, and magnetic) and dendrimers are used for imaging and targeted drug delivery [9]. Freitas Jr. [4] and Flynn and Wei [10] reported the importance of commercialization of nanomedicine including gated nanosieves, immunoisolation, ultrafast DNA sequencing, nanoshells, fullerenes-based pharmaceuticals, single-virus detectors, tecto-dendrimers, biologic robots and radio-controlled biomolecules [4]. This review deals with the practical application of nanomedicines and provides an overview of approaches in targeted drug delivery system.

Smart drug which are ideally designed to satisfy the requirements will pave the way for targeted drug delivery. These requirements includes: a) high stability (longevity) and not easily degraded or cleared by the reticuloendothelial system (RES) during blood circulation, b) accumulation of high concentration of drug at the tumor targeted sites or area, c) effective intracellular drug delivery at the tumour targeted area or site that matches drug pharmacodynamics of kinetics and spatial control and d) tolerability. The blood circulation stability and self-life of the drug (size, morphology, low degree of lipid hydrolysis and drug retention) are influenced by the physical and chemical nature of the encapsulated drug and encapsulation techniques. The design of tumor drug targeted delivery system based on: passive drug targeted delivery as a significance of enhanced permeability and retention effect (EPR) (without ligand targets and active drug release mechanisms) [11,12], targeted drug release based on recognition of receptor molecules [13-16], triggered drug release or stimuli – responsive release (pH, redox potential, temperature, light intensity, ROS response magnetic field, hypoxia response, glucose
and concentration of electrolytes) to localize the drug-nanocarrier to the determined targeted site [17-20].

**Physical and chemical interaction of nanocarrier surface with drug**

Tumor targeted drug should accumulate inside and/or outside of the nanocarriers by interaction with convalent and non-convalent linkage [21]. Encapsulation of targeted drug to nanocarrier is possible during the production of nanocarrier or after the carrier formation. Charging of therapeutic drugs is based on its solubility in the nanocarrier matrix, interaction of drug nanocarrier, the molecular mass of the therapeutic agent and the presence of functional group on a nanocarrier surface [22]. Nanocarrier molecules should hold high purity and chemical resistant nature which might play an important key role to control the targeted drug delivery. Nanocarriers are frequently altered (functionalization) before the encapsulation of targeted drugs [23]. Bioconjugation of polymer with therapeutic drugs leads to enhanced circulation time, minimal toxicity, biodegradation, controlled mechanical properties and well-organised drug release in the targeted sites and their characteristic nature might generate the barrier between cancer cells and nanocarriers [24-28]. Interaction of internal surface of the nanocarrier with entrapped drug molecules are protected beside early activation and biodegradation process therefore interaction of targeted drug with normal healthy cells is impossible [29,30]. Interaction of external surface of the nanocarrier with drug molecules is based on physical and chemical properties, bioconjugation and high surface area of nanocarrier molecules. Therapeutic drug molecule is interacting/bonding with functional groups of nanocarrier by covalently link to form amide, ester or acetyl-hydrazone and disulfide groups. Drugs with aromatic or plane structure can be adsorbed on the surface of nanocarrier by π-π interactions [31]. Furthermore, drug molecules accumulated on the external surface of the nanocarriers by electrostatic interactions (physical adsorption) [32].

**Bioconjugation between nanocarrier and targeted ligand**

Linkage between the targeted ligands with reactive groups of the nanocarriers surface can be categorised into covalent and non-covalent conjugation [33]. Common covalent linkage involve conjugation of two primary amines, two thiol groups, maleimide and thiol, a carboxylic acid and primary amine, hydrazide and aldehyde and a primary amine and aldehyde. Amide bond formation takes place in two stages: 1) triggering of carboxylic group on the surface of nanocarrier molecules, 2) activation of primary amine forming amide bond with nanocarrier molecules that enhance the activity of the applied amine forming amide bond with nanocarrier molecules. This methodology was discriminative and very effective on tumor targeted gene therapy. Thioether bond formed through the reaction between C1 carbon of maleimide and thiol group that is attached to the R, nanocarrier are stable within 24 hours in human serum even in the existence of reducing agent (e.g DTT) [33-35]. This reaction runs rapidly in aqueous solution as well as at room temperature. Inappropriately, the selectivity towards the thioether group formation is fairly low in aqueous solution, owing to side reactions such as formation of disulfides or intermolecular rearrangement. The thioether bond assures a longer duration for time distribution of drug and high discrimination of targeted drug delivery [34,36]. Alternative way is the coupling of thiol groups of the nanocarrier molecules with fragment of maleimide from the ligands these could be triggered via carboxylic acid or amine groups. This thioether and amide bonding are important in forming bioconjugation of nanocarrier-ligand system [36]. Conjugation of two thiol groups (originates from ligands and nanocarrier molecules) results in the formation of disulfide bond. Formation of disulfide bond between monoclonal antibody anti-My9 and liposomes acts strongly against HL-60 leukemia cells [37]. Bioconjugation of nanocarrier surface to the hydrazide group results in aldehyde groups of the ligands [33,34]. Basically ligands do not possess aldehyde structure but form through oxidation of hydroxyl groups. Conjugation of liposomes with antibodies C225 form acetyl-hydrazone group showed enhanced production of immunoliposome antibody in in vivo [38]. Advantage of this technique characterized by high immunological expression, prolonged distribution in the blood and can also easily control the immunoliposome synthesis. The cycloaddition reaction between a dienophile and a diene is called Diels-Alder (DA) reaction results in bicyclic compound formation. Coupling of nanocarriers and ligands enhances the yield (nearly to 100%) and can be synthesized easily under mild conditions [33]. DA reaction on ligands and nanocarrier linkages causes specific bond formation between the cancer cells and the ligand molecules [36]. Non-covalent bonding arranged by physical interaction of targeted ligands to the nanocarrier surface has an advantage owing to avoiding of severe destruction of therapeutic agents. Weak bonding, low control of reaction and binding of ligands in the undesired orientation are the weakness of the non-covalent interaction [39].

**Endocytosis of nanomedicines**

Drug targeted strategy are being validated to address important requirements for targeted therapy, drug delivery system and improved diagnosis. Tumor targeted drug delivery could be classified into carrier and non-carrier based therapy [40]. Carrier based drug therapies comprise, polymeric based drug carriers (polymeric nanoparticles [41], polymeric micelles [42], polymer-drug bioconjugates [43], dendrimers [44]), lipid based drug carriers (liposomes [45]), viral nanocarriers [46], inorganic nanocarriers (quantum dots [47], silica nanoparticles [48], carbon nanotubes [49]) and these nanomaterials with drug collectively called nanomedicine delivered the drug molecules, recombiant DNA and proteins to the targeted tumor cells/tissues/organs. Nanomedicines interact with surface of the targeted tumor cells and undertake intracellular signalling that determine the location (cytoplasm, nuclease) inside the targeted cells and improve the drug accumulation and acquire efficacy of targeted drug delivery [50]. The targeted drug should be stable in the blood circulation enhances the interaction with exterior of the plasma membrane which pave the way to targeted cells named as endocytosis [51,52]. Designed nanocarrier conjugated drug molecules enter into the tumor cells via endocytosis. The specific designed process of endocytosis consist of; a) particles recognition by opsonization in the blood circulation (complement components (C3, C4, C5); absorption of proteinks (immunoglobulins; IgG, IgM); and blood serum protein (albumin, fibronectin); b) attachment of opsonized particles into the membrane through receptor present on the targeted surface; c) ingestion of targeted drug by the cell; d) the mature phagosome fused with lysosome and form enzyme-rich phagolysosomes where
the nanomedicines prone to degradation and show its therapeutic indices [52,53]. Recent research proposed nanomedicine transport to the cell membranes based on essential characteristics nature of the nanocarrier (shape, size, hydrophobicity and surface charge) specific interaction between targeted ligand and cell membranes are designed as optimal targeted drug delivery system.

**Tumor targeted therapy**

Understanding of molecular mechanism important to tumor progression, formation, angiogenesis, invasion and metastasis clear and pave the way for drug development for targeted drug therapy based on cellular and molecular mechanism. Targeted drug alter the cell signalling pathways and molecular mechanisms (gene function) involved in the pathogenesis of diseases (cancer, parkinson etc...) offer an improved therapeutic value over conventional drugs. Tumor targeted therapy must satisfy few demands. Drug should be delivered to the malignant cells with minimum loss in the concentration and should selectively destroy the targeted tumor cells. Active form of drug release must be controlled simultaneously and minimal doses of therapeutic agents should be administrated for targeted drug therapy than the conventional therapy [54].

Targeted drug therapy also diminishes the possible side effects [36] owing the delivery of targeted drug molecules inside the cells and nanocarriers outside via the passive targeting mechanisms. The conventional therapy is linked with a drug aggregation process (form of carrier-drug) inside the malignant cells via EPR [55,56] thus, drug reaches based on the abnormal structure of the blood vessels near malignant cells [57]. Methotrexate [58], doxorubicin [57], paclitaxel [59], gemcitabine [60] and hexamethylmelamine [61] are most popular chemotherapeutic drugs used in conventional treatment. In tumor targeted therapy system ligands linked to the nanocarriers of therapeutic drugs facilitates the conjugation with specific receptor of the malignant cells. Thus, tumor targeted therapy supports the overexpression of tumor cell receptors have affinity of ligand towards receptor molecules [36,57,62] and delivery of therapeutic agents to the most resistant malignant cells and enhanced the prolonged circulation inside the tumor cells. This application provides high concentration of therapeutic drug inside the tumor cells and targeted drug cannot be released back to the blood circulation [55].

**Importance of nanoparticles as targeted drug delivery system**

The therapeutic value of drugs presently being used could be improved efficiently by delivering the drugs into the biological targets via nanoparticles. Drugs failed under the clinical trials might also be reconsidered using nanotechnological methods [63,64]. Drug concentration must be sufficient in the body to permit for an effective dose at the targeted tumor area. The targeted malignant cells must be strongly suppressed by its signalling mechanism essential for its cell survival with high toxicity towards the cancer or positive therapeutic strategies for nanomedicines. Using nanoparticles as carrier molecules a variety of targeted drugs including proteins, biological molecules, vaccines (viral and nonviral vectors), hydrophobic and hydrophilic drugs has been delivered [3]. Hence, eye-ball is a closed organ, an aptamer, antisense oligonucleotide for cytomegalovirus retinitis or a small interfering RNA (Si RNA) for age-related macular degeneration (AMD) being investigated in human eye [65]. Architecture of the nanostructured molecules are promising drug carrier that will assist the novel drug compounds towards the targets and has effects on

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**Figure 1:** Overview of targeted drug delivery system development from bench to bed.

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continuous drug discharge and capable of intracellular entry. Mostly nanoparticles [66,67] possess an advantages than microparticles that include: intravenous delivery, site-specific drug targeting for various diseases (human immunodeficiency virus infection, cancer, and central nervous system disorders), highly exploited for controlled drug release, prolonged circulation in the blood, facilitate extravasation and passive targeting, avoid opsonization owing to particle size less than 100nm and less concentration with prolonged action as well as enhancing the targeting of the drug to specific tumor sites [68]. Targeted drugs can also be administrated via transmucosal, pulmonary, transdermal and implantation delivery other than injection and oral routes.

Future Challenges and Direction

Application of nanomedicines and improvements of novel targeted drug delivery system play a significant role in tumor therapy. Nanocarriers minimize the enzymatic degradation of therapeutic agent and include consequently improve its activity. Furthermore, it enhances solubility, long-time bio-distribution of drug in the blood circulation and novel system break of biological barriers (blood brain) that improve the therapeutic drug delivered to the targeted tissues/cells/organisms. The targeted ligand nanocarrier molecules bioconjugated with drug must meets all basic requirements of safe and effective tumor therapy which include sufficient concentration, effective dosage, high cytotoxicity of therapeutic agents and aggregation of drug molecules on the external and internal surface of the nanocarriers might improves targeted drug delivery (Figure 1). Smart drug nanoparticles must be able to sense the exact alteration in their surroundings to target drug delivery. In addition to targeted drug delivery, smart drugs have application in regenerative medicines (self-regulating scaffolds for cell proliferation or infiltration or injectable systems for cell delivery) and tissue engineering. Development and synthesis of new polymeric based nanoparticles should be expected to be progressively elaborate and versatile nanocarriers of smart drug delivery for implementation in the future. Next generation of polymeric nanocarriers based drug delivery system with hormones, genes, growth factors, peptides, antibodies, etc… may enhance the efficacy and minimize side-effects. Therefore, improvement of multifunctional “smart” drug and nanocarriers pave the way for identification of cancer cells, drugs delivered to the targeted sites, real-time assessment of surgical and therapeutic efficacy, enhanced therapeutic index of multidrug resistance diseases and monitor the intracellular alteration to prevent precancerous cells from becoming malignant. Furthermore, in the future, merging expertise in the drug targeted delivery with technological improvements in nanomedicine should focus on ‘bench to bed-side’ practises to reduce delay to therapeutic stages.

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