Nanoparticles Mediated Drug Delivery

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Abstract

The term “nano” has originated from Greek word meaning extremely small or dwarf. Nanoparticles are nano-sized objects whose size is measured in nanometers (nm) ranging from 0.1-100 nm exhibiting distinct morphological characteristics which is quite different from their bulk form.

Keywords: Nano scale; Nano carriers; Quantum dot

Introduction

“Nanotechnology” the term was coined by Taniguchi [1]. It implies manipulation, reduction and fabrication of materials at nano scale with distinctive properties such as good strength, cost effective, lighter, ecofriendly, definite and specific etc and enhanced functionality and improved stability [2]. Nanotechnology has a wide range of applications; one of its potentials has been recognized in drug delivery systems for prevention and therapy of different diseases. Nanoparticles have been modified to effectively deliver drugs to the target regions in the human body. Besides, drug accumulation and its metabolism can also be tracked in the cells or tissues by using these tiny entities to get a better understanding of pharmacokinetics and pharmacodynamics of a particular drug [3].

Properties of Nanoparticles that Made Them Suitable for Drug Delivery

Nanoparticles being target specific have been used to deliver chemotherapeutics targeted to the tumor tissue without damaging normal organs [4,5].

Nanoparticles are chemically stable structures that could be fabricated through different chemical and biological routes that are environment friendly and cost effective.

These entities are designed in such a way that can easily diffuse into the cell by interacting with specific cellular components to permit selective targeting and accumulation in specific cell or tissues. Besides nanoparticles are pH-labile structures that easily degrades at low pH in the cell to release drug payloads. Examples of few biodegradable nanoparticles used for drug delivery system includes polymeric polyethylene-glycol etc. Nanoparticles can persist in the micro fluid environment of the cell for weeks or more without being degraded. Thus make them suitable for drug delivery systems. For example carbon nanotube can persist for a longer period of time with extensive functionalization and loading of cargo. Nanoparticles are designed in such a way that they are non-immunogenic i.e., compatible with the body immune system. As a result will not engage in initiating an immune response when introduced in the body [6].

Types of Nanoparticles for Drug Delivery

Numerous different types of nanostructures exhibiting different physiochemical properties are employed to improve the efficiency of drug delivery to specific targets [7,8]. Few examples are given below:

Polymersomes

Polymersome nanoparticles are synthetic amphiphilic blocks that are spherical vesicular bodies containing an aqueous solution. This unique copolymer is made of two components, a hydrophobic and hydrophilic subunits e.g. joined together. Poly lactic/glycolic acid, Polyethylene glycol-block-poly-caprolactone etc. [9,10].

Nanocapsule

Are also known as liposomes which are a lecithin and stearate encapsulated structure with a lipid core where lecithin is sited in the inner part of the capsule. Polyethylene glycol could be employed to increase its half-life [11,12].

Silica nanoparticles

These are mesoporous nanoparticles with high specificity in terms of pore volume and surface area. Differential surface enhancements are employed to optimize particle size which directly influences its function. Thus making them suitable to be used in imaging tools [9].

Quantum dot

Amalgamated nanoparticles composed of two subunits Poly L-lactic acid-block-polyethylene glycol. High optical resolution, small size and property of fluorochrome made them useful for tracing the dynamics of many biological molecular pathways [13]

Amphiphilic nanoparticles

These dual nature nanoparticles consisted of hydrophobic and hydrophilic regions. Hydrophobic regions provide protection against the polar surroundings forming a micelle. In case of drug delivery it carries the drug and provides protection from the body’s immune system to prevent its elimination e.g. polyethylene oxide-polyaspartic acid copolymers micelle carrying the drug doxorubicin [14].
Dendrimers
Repetitively branching nanostructures with highly specific large surface area making them target specific. It is mostly composed of polyamidoamines [15,16]. For example telodendrimers made of polyethylene glycol and dendritic cholic acid subunits [17,18].

Graphene
Allotropic form of carbon with unique thermochemical properties and high tensile strength. Its unique hexagonal structure enables infrared radiation conversion in to heat. High surface area allows drug delivery in cancerous cell but is quite toxic in nature therefore requires surface shielding [19,20].

Carbon nanotubes
These are basically graphene nanotubes used as hybrid drug carries. When bombarded with near-infrared radiation, it allows controlled drug release at specific tumor sites to protect the neighboring normal tissues [21-23].

Metal-core nanoparticles
Such nano entities are magnetic and photothermally stable. In response to an external magnetic field these entities vacillates and heat up e.g. Magnetite and maghemite [24].

Oligopeptides
pH sensitivity zwitterionic oligopeptide lipids nanoparticles mostly used as nanocarriers [25].

Supramolecularpolymers
Due to their reversible monomer-to-polymer transitions efficient in drug delivery [26].

Drug Delivery Targeting
There are three main methods to transport drug-loaded nanoparticles to diseased sites:

Passive targeting
Enhanced permeation and retention effect is the mechanism behind passive targeting which is the property of tumor cells to absorb particles of specific size than normal cells due to drippy tumor blood vessels and faulty particle screening [27]. Nanoparticle morphology and electrochemical properties influence enhanced permeation and retention effect thus effecting drug penetration, duration of circulation and intracellular stability [28,29].

Active targeting
Active targeting involves the use of ligands such as antibodies, proteins, and peptides bound to nanoparticle surface to increase their uptake selectivity by interacting with the overexpressed receptors at the target sites [30]. These ligands often protect nanoparticles from enzyme destruction. High binding affinity of ligands to the target cell will increase drug delivery efficiency [31].

Physical targeting
Physical targeting uses external sources or fields to guide the nanoparticle to the target site and also controls the release process, for example, in photothermal and magnetic hyperthermia therapy [32].

Challenges in nanoparticle designing for drug delivery
Many challenges have been encountered when designing medically significant nanoparticles for drug delivery. The classified and nonuniform nature of nanoparticles is a prime challenge encountered during nanoparticle designing. A small alteration in a single existing property can have deleterious consequences on the pharmacokinetics and pharmacodynamics of the particle [33]. For example maintaining a narrow distribution of particle size is an issue. Particle size under 200 nm is desirable but with a broad normal distribution it is difficult to limit the number of particles over 200 nm [34]. In addition, variation in nanoparticle structure must be smaller otherwise the designed nanoparticle works in a lab setting will not work in an actual cellular environment. Nanoparticle stability is one of the major concerns that it should be stable in solution and should not aggregate over time. Finally, nanoparticles toxicity should be minimized because it greatly increases the time and cost of clinical trials. These challenges must be overcome in order to frame nano entities with enhanced properties thus making them suitable for drug delivery [35].

Conclusion
Nanoparticles are excellent candidates for targeted drug delivery. Being stably biocompatible and readily available allows for precise doses of drugs to be encapsulated and delivered directly to the target site. However nanoparticles morphology, electrochemical and thermal properties should be controlled to avoid side effects. Environmental factors can also impact nanoparticle drug delivery efficacy. Therefore a nanoparticle aimed for drug delivery must undergo clinical trials before practical application to ensure that they do not cause harm to living tissue or animals.

References


