Nanomedicines as Cancer Therapeutics: Current Status

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Abstract: As of 21st century, cancer is arguably the most complex and challenging disease known to mankind and an inevitable public health concern of this millennium. Nanotechnology, suitably amalgamated with cancer research, has ushered an era of highly personalized and safer medicines which can improve cancer diagnosis and therapy. A wide variety of nanomedicines are currently under investigation, including polymeric/non-polymeric nanoparticles, dendrimers, quantum dots, carbon nanotubes, lipid- and micelle-based nanoparticles. The bases of these nanomedicines in reducing toxicity associated with cancer therapy are their ability to carry a large payload and multivalent-ligand targeting. This imparts specificity for targeting the tissues as well as bypass resistance mechanisms. The major hurdles on these future medicines are potential toxicity of nanoparticles, which imposes the need of extensive regulatory evaluation before nanomedicines could be utilized as cancer therapeutics. This review highlights nanopharmaceuticals that have been investigated in oncology for various applications (diagnosis, therapeutic delivery and theranostics). It also discusses the effects of nano-sized materials on tissues/organ functions, the possibility of overcoming multi-drug resistance by using nanomedicines and their current clinical status.

Keywords: Cancer nanomedicine, multi-drug resistance, nanotechnology, nanotoxicity, regulatory, theranostics.

INTRODUCTION

The practice of cancer therapy has changed considerably in the past decades. Curative treatments have been achieved for a number of fatal malignancies such as testicular cancer, lymphomas and leukemias. However, there are still numerous constraints associated with the use of conventional chemotherapeutic agents. Although some of the newer agents indirectly target the survival of tumor cells, viz. bevacizumab (Avastin) inhibits tumor angiogenesis [1], and ipilimumab is an immunotherapeutic antibody [2], the majority of them act by directly killing the cells in a more or less specific manner. Such agents are associated with dose-limiting side-effects and anticancer drug resistance, since complete elimination of malignant cells is often not possible using therapeutic doses. To overcome these problems, various approaches are being developed that use nanocarriers for targeting tumor site via passive or active means [1, 2]. According to the US Food and Drug Administration (FDA), nanotechnology product refers to products that are at least one dimension in the length scale between 1-100 nm and display functional behavior in relation to their nanosized properties [3]. Tailor-made nanomaterials can be useful for reducing multi-drug resistance, evading blood-brain barrier (BBB), and diagnosis and imaging of cancer [3]. Even though the research in pharmaceutical nanotechnology is in a very active stage, approved nanomedicines in the clinical practice is very low. However, it is encouraging to observe that there are many nanoformulations that are either approved, or in different phases of clinical trials [4]. The commercially available nanomedicines that are clinically employed in cancer treatment are Doxil®/Caelyx® and Abraxane®. Doxil® is a liposomal formulation of doxorubicin, which is approved for treatment of Kaposi’s sarcoma, refractory breast and ovarian cancer. Abraxane® is an albumin-based formulation of paclitaxel that has been approved for the treatment of metastatic breast cancer. Despite the progress of some nanomedicines in clinical practice, there are still issues related to their tissues/organ biodistribution, toxicity, environmental and regulatory aspects. This manuscript reviews mechanisms of action, toxicity, and regulatory aspects, recent advances, and future prospects of different nano-carriers developed as anticancer therapeutic and diagnostic agents (theranostics).

TARGETING STRATEGIES AND ROLE OF NANOMEDICINE

Tumor blood vessels possess distinctive pathophysiological characteristics that are not observed in normal blood vessels, viz. a relatively high proportion of proliferating endothelial cells, increased tortuosity, pericyte deficiency and aberrant
basement membrane. Moreover, tumors also feature a decreased lymphatic drainage resulting in increased retention of extravasated macromolecules. This phenomenon is described as “enhanced permeation and retention (EPR) effect” [5-7], which is often proposed as the underlying mechanism for the accumulation of nanocarriers in tumors, and forms the basis of passive targeting. Alternatively, nanocarriers can accumulate in tumors by active targeting strategy, which involves binding of nanomaterials to surface-exposed receptors on tumor cells or tumor blood vessels. Passive and active targeting will be discussed in more detail in the later section. Therefore, irregular tumor vasculature and impaired lymphatic drainage create high interstitial fluid pressure with an extremely hydrophilic environment [6], which may limit the distribution of drugs to solid tumors. Likewise, the presence of extracellular matrix of the tumor, fibrillar collagen, and necrotic non-supporting regions serve as barriers for the effective delivery of drugs [6-9]. It has been proposed that antiangiogenic agents may normalize an efficient blood flow to tumors, which will decrease the hypertensive interstitial condition, thereby improving the delivery of chemotherapeutics [10]. Drugs that have been shown to successfully normalize the tumor vasculature include anti-VEGF antibody bevacizumab [11].

Passive Targeting

Fast growing vascularization, leakiness and defective lymphatic drainage all contribute to the retention of macromolecules and nanoparticles in tumors, which is the rationale for EPR-based drug targeting. Studies with liposomes and other nanoparticles have indicated that the cut-off size of the pores in tumor vessels is as large as 200 nm–1.2 μm [12-15]. In fact, it is now established that nanoparticles with a size range of 10–100 nm can accumulate in tumor tissue via EPR effect [12-13]. Both, the size of nanoparticles and their surface properties must be tailored appropriately to avoid phagocytosis by the RES (reticuloendothelial system). Hydrophilic surface coating of nanoparticulate systems by PEG, poloxamers, poloxamine, polysaccharide and branched/block amphiphilic co-polymers have been successfully applied for this purpose [12-14].

Active Targeting

Active targeting approach is based on the interaction of a ligand-equipped nanocarrier with surface-exposed receptors on target cells, which will help in their accumulation in tumor and, more importantly, will facilitate their intracellular accumulation via receptor-mediated endocytosis [15-17]. Most cancerous cells have one or more types of over-expressed molecular targets which may serve as the site for active targeting through nanomedicine. Endocytosed particles are transported to endosomes and subsequently to lysosomes, where further processing of the particles and release of drug(s) is affected by the presence of lysosomal enzymes and low pH of this intracellular compartment. Active targeting mechanisms provide an alternative means to combat multi-drug resistance (MDR) because resistance inducing proteins such as P-glycoprotein (P-gp) cannot pump out nanoparticle-associated drug or drug-polymer conjugates that have entered the cell via the endocytic mechanism [15-21].

Folate Receptor–Based Targeting

Folic acid, an oxidized form of folate, is a vitamin required for one-carbon transfer reactions and is essential for the de novo biosynthesis of DNA and several nucleotide bases [22]. Folate transport across plasma membrane in normal cells is carried out either by folate carrier or by folate receptor [23]. Folate receptor is a highly selective tumor marker, which is overexpressed (100- to 300-times higher than in normal tissue) in many human cancer cells such as brain, kidney, breast and lung [24]. Therefore, folate-based targeting ligands find potential applications in the diagnosis and delivery of chemotherapeutic agents [23]. Their high receptor affinity and restricted interaction with normal tissues facilitate the delivery of therapeutic agents selectively to the target site [25-28].

Monoclonal Antibody–Based Targeting

Monoclonal antibodies (Mabs) are antibodies obtained from a single clone of a parent immune cell. These are immunoglobulins based on complex proteins, which contain mainly two regions; the antigen-binding fragment (Fab) and complement fixing fragment (Fc), which are responsible for specific antigen binding and fixing complement for in-vivo biological response [29]. Primary tumors as well as metastatic tumors generally over-express certain antigens on their surfaces (like HER2 in breast cancer, epidermal growth factor receptor in lymphoma, etc.). Mabs specific to particular antigens provide an ideal platform for conjugated drug targeting [30]. Current research is being focused on the development of chimeric and fully humanized derivatives of such targeting Mabs to decrease their immunogenicity [31]. PLGA immuno-nanoparticles conjugated with Mabs directed to breast cancer cytokeatins interacted specifically with MCF-10A neoT [32]. Steinhauser and co-workers suggested trastuzumab-modified nanoparticles for the treatment of HER2-over-expressing breast cancer cells [33]. In 2004, Gemtuzumab-ozogamicin (Mylotarg), which is targeted to the cell-surface receptor CD33, was approved by FDA for the treatment of acute myelogenous leukemia (AML) [34]. This conjugate was however withdrawn from the market in 2010 due to fatal toxicity in phase-3 clinical trials.

Beside this, several other drug-antibody conjugates have been developed. Some of them are already approved while others are in different phases of clinical trials. From this category, Brentuximab vedotin that binds to CD30, a tumor-specific marker of the TNF-receptor superfamily, has recently been approved for the treatment of Hodgkin and systemic anaplastic large cell lymphomas [35, 36]. Additionally, Inotuzumab ozogamicin (target: CD22) and Trastuzumab emtansine (target: HER2) have shown pronounced clinical activity in phase 3 clinical trials for non-Hodgkin lymphoma, acute lymphocytic leukemia and metastatic breast cancer [35, 37].

CD95 Targeting

CD95 is a tumor necrosis factor (TNF) receptor that prompts cell apoptosis of peripheral T-cells upon oligomerization [38]. CD95-mediated immunological destruction of liver tissue (in hepatitis) and the CD95 death system’s role in killing infected hepatocytes is well established.
McCarron et al. reported a novel immuno-nanoparticle against colorectal tumor cells. Nanoparticles were peripherally coated with antibodies directed to CD95 and loaded with camptothecin. Effective internalization was confirmed by fluorescence visualization studies in HCT116 cell line. Similarly, they developed camptothecin nanoparticles without anti-CD95 Mab, which showed poor internalization kinetics [39].

**Transmembrane Tyrosine Kinase Receptor Targeting**

Huh et al. successfully worked on metallic nanoparticles tagged with trastuzumab leading to localized intratumoral accumulation *in-vivo* [40]. Akira et al. encapsulated metallic nanoparticles (MNPs) in anti-HER2 immunoliposomes and used these antibody-directed nanoparticles for hyperthermia therapy [41]. Similarly, gold nanoshells were developed for targeting and photothermal therapy of HER2 over-expressing ovarian cancer OVCAR3 cells. OVCAR3 cells that had internalized the metallic nanoparticles were irradiated with near infra-red (NIR) radiation leading to selective destruction of cancer cells through photothermal ablation [42]. Parallel to this theme, Day et al. developed gold-gold-sulphide nanoparticles that were used in photo-ablation therapy. These nanoparticles could be imaged via multiphoton microscopy and could induce thermal damage via hyperthermic NIR [43].

**Aptamer-Based Targeting**

Identification of target proteins having a clear link with human oncologic disorders offers a powerful tool for the development of relevant aptamer ligands. Aptamers, a class of macromolecules containing single-stranded RNA or DNA (ssRNA or ssDNA), can be selected from random pools, based on their antigen binding affinity [44, 45]. A group of researchers have hypothesized an automated selection process (termed as “systematic evolution of ligands”) by exponential enrichment for rapid isolation of RNA ligands and aptamers [46]. Aptamers possess many favorable characteristics such as ease of chemical isolation, selective binding affinity, small physical size, and lack of immunogenicity, which make them useful as homing ligands in nanomedicine-based cancer therapy [47-50]. Furthermore, aptamers can be surface-modified with various functional groups to facilitate their conjugation with nanomaterials, which could be exploited for their selective targeting in the diagnosis and therapy of cancer [46, 48, 49]. Nanoparticles (PLGA–block-PEG copolymer) along with A10 aptamer bioconjugates have been targeted to prostate-specific membrane antigen (PSMA), which is a transmembrane protein responsible for prostate cancer [49]. In another study, a marked reduction in tumor size was observed over a period of 109 days after a single intra-tumoral injection of aptamer-decorated nanoparticles [50]. Their specific binding affinity with PSMA cells was further confirmed by *in-vivo* cellular imaging techniques using an aptamer conjugated to luminescent CdSe and CdTe nanocrystals, which were found to be more concentrated in prostate tumor cells. Research is being directed for the development of aptamers that can bind selectively to vascular endothelial growth factor (VEGF), a protein involved in angiogenesis. They would especially be useful in the treatment of age-related macular degeneration [51].

**Integrin-Based Targeting**

Integrins were originally characterized as a family of cell adhesion receptors that bind to extracellular matrix and cell-surface ligands. The role of cell-surface integrins is to transmit and detect changes from the extracellular matrix to intracellular signaling, and thus regulate tumor growth, angiogenesis, proliferation, apoptosis and metastasis [52]. Several integrins play a key role in promoting tumor angiogenesis and metastasis. Within the family, αβ3 integrin adhesion receptor is the main marker of angiogenic blood vessels in the mammalian system, and is over-expressed on growth factor–activated endothelial cells. A synthetic peptide bearing Arg-Gly-Asp sequence specifically binds to the αβ3 integrin and can be used to selectively target angiogenic blood vessels [53].

**Drug Like Peptides: Vasoactive Intestinal Peptide**

Vasoactive intestinal peptide (VIP) is a 28-amino acid neuropeptide of the glucagon-secretin category that is distributed widely in both the central and peripheral nervous systems. VIP receptors are five-times more abundant in breast cancer cells than in normal breast cells. Additionally, VIP is synthesized locally within the eye and takes part in immunological homeostasis of the ocular microenvironment. However, its potential therapeutic applications are restricted because of its failure to cross the blood-brain barrier (BBB), rapid elimination and degradation (half-life <1 min) after i.v. administration. Therefore, various formulation-based approaches have been explored to protect its degradation and make it available in the intact form, mostly via the nose-to-brain pathway. For brain delivery, a VIP analogue was conjugated to OX26 monoclonal murine antibody and directed against the rat transferrin receptors [54]. Embedding VIP into wheat germ agglutinin–coated PEG-PLA or protamine-oligonucleotide nanoparticles enhanced their transport to the brain after intranasal administration [55]. The VIP receptor, in addition to HER-2 and estrogen receptors, is present in high density and considered as the biomarker or molecular target for breast cancer. The overexpression of VIP receptors is increasingly being used to develop new drugs for the treatment of cancer patients. Targeting PEGylated liposomes, which contain radioisotopes to VIP receptors of tumor, resulted in an enhanced passive and active breast cancer inhibition in rats [56].

**Luteinizing Hormone-Releasing Hormone (LHRH) Targeting**

Luteinizing hormone-releasing hormone (LHRH) is another targeting moiety. The LHRH receptor is barely present on the surface of most healthy human cells, but is over-expressed in ovarian and some other cancer cells [57, 58]. Dharap et al. recently developed the LHRH–PEG–camptothecin targeted anticancer drug delivery system, wherein LHRH targets the corresponding receptors in cancer cells, PEG prolongs the blood circulation time, and camptothecin functions as the anticancer drug [58]. The targeted conjugate exhibited significantly higher cytotoxicity against cancer cells than the non-targeted PEG–camptothecin conjugate or the free drug *in vivo*, indicating the validity of actively targeted nanoparticles for anticancer therapy. The
list of various targeting ligands and their respective targets are given in Table 1.

CLASSIFICATION OF NANOCARRIERS IN ONCOLOGY

Advancement and growing interest in nanotechnology-based drug delivery systems and imaging agents have dramatically increased their market potential. The market of nanopharmaceuticals is growing by 17% and expected to have a $53 billion market in 2014. Hence, a steady succession of nanomedicines is anticipated to seek regulatory approvals and subsequent access to human use. Nanomedicines have several features that make them especially adept at carrying anticancer compounds to tumors. First, they are large enough to avoid rapid elimination through the kidney, yet they are small enough that they may penetrate the leaky vasculature of the tumor tissues and get trapped because of the EPR effect; nanoparticles only show this EPR effect in tumors or inflamed tissues that also have a leaky vasculature, while they avoid normal tissues to a large extent since they cannot penetrate the intact endothelial barrier. Second, nanomedicines can enter a cell by endocytosis very efficiently especially when decorated with targeting ligands. Furthermore, after intracellular accumulation, nanocarriers protect the particle’s payload from being expelled by cellular pumps that otherwise are responsible for drug resistance. Nanoparticles are useful carriers especially for drugs that operate intracellularly such as interference RNA. Another attractive feature of nanoparticulate systems is that they can incorporate multiple drugs, making combination therapy possible on a single platform. On the basis of application in oncology, nanopharmaceuticals may be classified as therapeutics, diagnostics or theranostics (i.e. combination of both). The developed nanocarriers for medicines are illustrated in Fig. (1). Diverse classes of nanomedicines have been developed till now and they have their own advantages and disadvantage as drug carriers (Table 2).

Liposomes

Liposomes are a family of microscopic, concentric lamellar structures enclosing an aqueous volume by a membranous lipid bilayer. They are formed by blending natural or synthetic phospholipids, cholesterol and tocopheryl acetate, and their subsequent hydration in the aqueous media. Liposomes have been studied and evaluated extensively for the spatial and temporal delivery of anticancer agents. First liposomal product approved by FDA is Doxil® in 1995 for the treatment of Kaposi’s sarcoma, refractory breast and ovarian cancer. Liposomes qualify as suitable candidates for delivering drugs in oncological disorders owing to their straightforward formulation methodology, as well as their versatility in entrapping both hydrophilic and lipophilic drugs [59-61]. PEGylated or stealth liposomes (long circulating) exhibit remarkable inhibition of the rapid uptake by the RES cells, leading to an improved extravasation profile and enhanced intratumoral localization. Conjugation of a targeting ligand to liposomes can take the tumor targeting approach to a higher level. A study conducted pertaining to the uptake of folate-PEG-

<table>
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<tr>
<th>Targeting Ligands</th>
<th>Specific Targets</th>
<th>Location</th>
<th>Targeting Application in Oncology</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>NGR peptide</td>
<td>Aminopeptidase N</td>
<td>Cell surface</td>
<td>Vasculature in solid tumors</td>
<td>[150]</td>
</tr>
<tr>
<td>Ab &amp; Ab-derived peptides</td>
<td>Aminopeptidase P</td>
<td>Caveoli</td>
<td>Lung cancer</td>
<td>[151]</td>
</tr>
<tr>
<td>Antibody (Rituxan, Zevalin)</td>
<td>B-lymphocyte antigen CD20</td>
<td>Cell membrane surface</td>
<td>B cell lymphomas</td>
<td>[152, 153]</td>
</tr>
<tr>
<td>Ab (Erbitux) aptamers</td>
<td>EGF receptor (ErbB1)</td>
<td>Cell membrane surface &amp; lipid rafts</td>
<td>Metastatic colorectal cancer</td>
<td>[154]</td>
</tr>
<tr>
<td>Ab, Ab-derived peptide</td>
<td>Endoglin</td>
<td>Cell membrane surface and caveoli</td>
<td>Vasculature in solid tumors</td>
<td>[155]</td>
</tr>
<tr>
<td>Peptides, Ab, Ab-derived peptides</td>
<td>ICAM-1</td>
<td>Cell membrane surface</td>
<td>Vasculature in solid tumors</td>
<td>[156]</td>
</tr>
<tr>
<td>Ab (Avastin), peptides</td>
<td>VEGF receptor</td>
<td>Cell membrane surface</td>
<td>Vasculature in solid tumors</td>
<td>[157]</td>
</tr>
<tr>
<td>Ab &amp; Ab-derived peptides</td>
<td>PECAM-1</td>
<td>Cell membrane surface</td>
<td>Lymphoid cancers</td>
<td>[158]</td>
</tr>
<tr>
<td>Ab aptamers</td>
<td>MUC1</td>
<td>Cell membrane surface</td>
<td>Breast &amp; bladder cancer</td>
<td>[159]</td>
</tr>
<tr>
<td>Albumin &amp; Ab</td>
<td>gp60</td>
<td>Caveoli and Cell membrane surface</td>
<td>Vascular targeting in malignant liver cancer</td>
<td>[160]</td>
</tr>
<tr>
<td>Trastuzumab (Herceptin), pertuzumab Ab, tyrosine kinase inhibitors</td>
<td>EGF (ErbB) receptor (ErbB1) and EphA2 receptor</td>
<td>Cell membrane surface and lipid rafts</td>
<td>Breast, prostate cancer, and Metastatic colorectal cancer</td>
<td>[161]</td>
</tr>
<tr>
<td>Ab, Ab-derived peptides, aptamers</td>
<td>insulin-like growth factor receptor (IGF-1R)</td>
<td>Caveoli and Cell membrane surface</td>
<td>Lungs, pancreas and breast cancer</td>
<td>[162, 163]</td>
</tr>
</tbody>
</table>
liposomal doxorubicin by KB cells (a cell line derived from human carcinoma of nasopharynx) showed 45-fold higher uptake of folate-targeted liposomes than of liposomal doxorubicin and 1.6-fold higher uptake versus free doxorubicin, while cytotoxicity was 86 and 2.7 times higher, respectively [62]. Opsonisation of plasma proteins to nanocarriers greatly enhances their clearance. To diminish this, albumin was conjugated to PEGylated doxorubicin liposomes [63]. The disposition of this type of nanoparticulate doxorubicin to the heart was significantly smaller than that of free drug, and on the other hand tumor accumulation of albumin-conjugated PEGylated liposomes was higher than the PEGylated liposomes. Liposomes can also be used to overcome the specific resistance mechanisms such as P-gp mediated drug efflux, and targeting drugs to tumor tissue and cells, leading to enhanced bioavailability at the site of action to the resistant tumors safely and effectively [64]. Currently, liposomal anthracyclines have achieved clinical use in cancer treatment by eliciting enhanced encapsulation of drug in a stable, non-reactive carrier, providing potential benefits in case of resistant tumors.

Niosomes

Nonionic surfactant-based vesicles (NSVs) are now one of the most widely studied alternatives to liposomes, resulting from the self-assembly of hydrated surfactants monomers. Surfactants can be employed as useful surrogates to phospholipids in the fabrication of vesicular systems [65, 66], and have some advantages over liposomes like avoidance of superfluous use of phospholipid (vulnerable to oxidative degradation) thereby boosting formulation stability, and drastic reduction in production cost. The rationale behind using niosomes in anticancer therapy is to prolong the circulation of entrapped drug, altering its organ distribution and metabolic stability, as well as to reduce drug-induced toxic effects to extra-tumorous tissues. Like liposome, PEGylation and surface modification for enhanced tumor targeting is also possible. Lately, it has been shown that efficient tumor targeting of hydroxycamptothecin-loaded pegylated niosomes is possible when modified with transferrin [67].

Polymeric Micelles

The promising potential of polymeric micelles (PMs) as effective drug delivery carriers in anticancer therapy has been a topic of discussion since decades. The major breakthrough was witnessed in the early 1990s by Kataoka’s group, who developed doxorubicin-conjugated block copolymer micelles [68, 69]. Since then, a large number of modified pendant polymeric micelles have been used for the delivery of chemotherapeutic agents in pre-clinical and clinical studies. A polymeric micelle essentially consists of an inner hydrophobic core (which is the part of block copolymer) and encapsulates a hydrophobic/poorly-water soluble drug, while the outer shell or the corona is composed of the hydrophilic part of the block copolymer. Some inherent properties of PMs which make them an obvious choice for carriers of anticancer drugs are: (1) size in nano range; (2) stability in plasma; (3) in vivo longevity owing to hydrophilic corona as observed in PEGylated (stealth) liposomes; and (4) EPR effect by virtue of the pathological characteristics of the tumor cells, is observed allowing passive targeting to the tumor cells [70]. The corona serves the dual purpose of stabilizing the PMs against recognition by RES cells and targeting the drugs by attaching specific ligands recognizing the tumor sites. Drug release can be coordinated and regulated by appropriate application of heat or ultrasound, or other external suitable stimuli. Thus, it can be conveniently accepted that PMs have the potential of active as well as passive targeting to the tumor tissues.
Table 2. Common preparation methods, advantages and limitations of different nanocarriers in anticancer drug delivery.

<table>
<thead>
<tr>
<th>Nanomedicines</th>
<th>General Preparation Technique</th>
<th>Advantages</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polymeric nanoparticles</td>
<td>In-situ polymerization, nanoprecipitation, supercritical fluid technology, gelification, emulsion-solvent evaporation, Interfacial polymerization of monomers or phase inversion process with emulsions of polymers</td>
<td>Multiple drug loading, Controlled release can be achieved for even more than 1 month by tailoring the mechanical properties and selection of polymer</td>
<td>Stability and re-dispersability not always acceptable, presence of residual solvents, sometime burst release</td>
</tr>
<tr>
<td>Solid lipid NPs</td>
<td>Ultrasonication/high speed homogenization, SLN preparation by using supercritical fluid, High-pressure homogenization, dilution of microemulsions or solvent removal from oil-in-water emulsions, solvent diffusion technique and freezing of an emulsion of lipids</td>
<td>Established Biocompatible, Flexibility of size and surface manipulation, lower toxicity vs non-liposomal formulation, Feasibilities of carrying both lipophilic and hydrophilic drugs, Water based technology (avoid organic solvents), Easier to validate and gain regulatory approval</td>
<td>Poor drug loading capacity, drug expulsion after polymeric transition during storage and relatively high water content of the dispersions, Poor batch to batch reproducibility Sterilization difficulties, Release of drug not always well controlled</td>
</tr>
<tr>
<td>Liposomes and Niosomes</td>
<td>Film hydration method, Reserves Phase Evaporation Method, Freeze-Thaw Method, Extrusion, Solvent Injection Methods and Sanitation Method</td>
<td>Liposome particularly made up of natural lipids, Encapsulate both hydrophobic and hydrophilic drugs, surface functionalization is easy, Lower toxicity particularly in case of case of liposome</td>
<td>Fair stability Poor batch-to-batch reproducibility Difficulties in sterilization Low drug loading</td>
</tr>
<tr>
<td>Polymeric micelles</td>
<td>Direct or controlled aggregation of block copolymers in a solvent, direct dissolution, solvent evaporation, and dialysis</td>
<td>Long circulation time, effective EPR, Effective for hydrophobic drugs, good safety profile, Very small size (10-100nm), good leading capability, helpful for MDR, Simple functionalization for the targeting, easy of sterilization</td>
<td>Difficult polymer synthesis, Not good for hydrophilic molecules, In-vivo instability, Slow extravasation, risk of chronic liver toxicity due to slow metabolic process</td>
</tr>
<tr>
<td>Dendrimers</td>
<td>Convergent and divergent synthesis method of chemical polymerization</td>
<td>Formulation having particle size range from 1 to 100 nm, surface functionalization is easy due to presence of multiple functional groups</td>
<td>Monotonous preparation techniques and toxicity is also concern because of reaction solvent.</td>
</tr>
<tr>
<td>Metallic nanoparticles/magnetic nanoparticles</td>
<td>Chemical reaction (Photoreduction), Laser ablation, Sonochemical reactions, Low-temperature vapour co-condensation,biological method</td>
<td>Availability of variety of metallic nanoparticle, theranostic carrier, targeting is easy due to presence of multivalent surface, multiple payload, easy to achieve small size even of 5nm, excellent EPR</td>
<td>Stability and control chemical reaction is challenging work, toxicity due to reactive surface</td>
</tr>
</tbody>
</table>

Tumor angiogenesis and tumor cell targeting of PMs is a prospective application [71].

**Polymeric Nanoparticles**

Encapsulation, dissolution, entrapment of drug in nanosized particles or their attachment to a polymeric matrix gives rise to polymeric nanoparticles (NPs). A plethora of well-studied biodegradable polymers like PLA, PLGA, Poly-ε-caprolactone, etc., are used to fabricate polymeric NPs, maximizing tissue compatibility and minimizing toxicity due to their biocompatibility and biodegradability. Alternatively, synthetic polymeric agents can also be used to formulate NPs and the accompanied advantages include increased biological half-life, reduced uptake by the RES cells and improved targeting to tumor cells. Mu and Feng synthesized PLGA NPs containing the drug Paclitaxel (PTX) (Taxol®) for solubility enhancement [72]. Gryparis and co-workers observed the *in vitro* anticancer activity of cisplatin-loaded PLGA-mPEG NPs on human prostate cancer LNCaP cells. It was found that as the PLGA/PEG ratio in PLGA-mPEG copolymer increased, the cytotoxicity also increased. This effect was observed primarily due to increased uptake by cells with the increasing ratio. Furthermore, it was also concluded that the *in vitro* activity of cisplatin was comparable to free cisplatin [73]. Li et al. prepared multifunctional pluronic/polyethylenimine NPs conjugated with folic acid (as folate) to their surface to improve targeting to tumor cells. They concluded that the surface-modified PTX nanoparticles showed sustained release profile in contrast to PTX solution [74]. Additionally, the *in vitro* anticancer activity was also enhanced considerably as compared to free drug, and folate conjugated nanoparticles demonstrated excellent cytotoxic action. In a similar experiment conducted by Mo and Lim, PTX-loaded PLGA NPs were conjugated with wheat-germ agglutinin (WIT-NP) which showed superior cytotoxicity towards A549 and H1299 cell lines as compared to conventional PTX formulations (as measured by IC-50 doses). This superiority was attributed to a more efficient and complete intracellular
PTX accumulation via WGA-receptor mediated endocytosis and IPM-facilitated intracellular drug release. WIT-NP cytotoxicity can be ascribed to PTX-induced tumor cell apoptosis and cell cycle arrest at G2/M-phase [75].

**Solid Lipid Nanoparticles (SLN)**

SLNs offer an attractive and efficient means of drug delivery, particularly for hydrophobic, poorly water soluble drugs. They combine the advantages offered by liposomes, NPs and fatty emulsions. SLNs are comparatively stable colloidal carrier systems in which the drug is entrapped in a lipid core, and manufactured by high-pressure homogenization or microemulsification technique [76]. Significant advantages presented by SLNs over polymeric systems are their low toxicity due to lipid biodegradability, and small size which helps to bypass the RES.

**Inorganic Nanoparticles**

The synthesis of inorganic NPs is an active area of application research in nanotechnology. Some common metals of utility in anticancer drug delivery include gold (Au), copper (Cu), and silver (Ag), and other non-metallic inorganic carriers, such as silica. Among these, gold NPs have emerged as superior carriers following their excellent optical and photoelectric properties, inertness and non-toxicity, higher stability, ease of preparation, scope for bioconjugation and bio-modification with thiol, disulfides and amino groups. Their dispersibility can be augmented by conjugation with thiolated-PEG [77]. Important areas of gold NPs applications are their use as contrast agents in diagnosis, and in photothermal cancer therapy. In recent years, gold NPs are most extensively evaluated in oncology as compared to the other metallic NPs. Gold NPs have unique properties that make them potential candidates in therapeutic oncology [78-81]. Gold nanomaterials have optical properties with high absorption efficiency particularly in the IR region that can easily be enhanced by changing the shape and surface area at the nano scale [81]. Gold nanomaterials in particular shapes like nano-rods, nano-stars, nano-cages and nano-shells show excellent localized surface plasmon resonant properties which particularly favor their applicability in oncology. The highly tunable and multivalent surface of gold NPs favor high payload by covalent or non-covalent conjugation [82, 83]. Furthermore, it is possible to functionalize the gold NPs for active targeting with targeted ligands. For example, PEGylated gold NPs have been conjugated with human transferrin (Tf) by amine-carboxylate reaction that showed significantly improved uptake of particles in cancer cells after i.v. administration to mice bearing Neuro2A tumors [84].

Shen and co-workers demonstrated effects of functionalized gold NPs on the binding of dacarbazine to DNA bases. They illustrated the apparent enhancement of the electrochemical response of dacarbazine revealing the facilitation of specific interactions between the drug and DNA bases [85]. Cherukuri et al. presented an informative review on the emerging role of NPs especially gold in the hyperthermic treatment of cancer. Metallic NPs induce hyperthermic cytotoxicity when exposed to near-infrared radiation or radiofrequency fields [86]. Recently, a couple of reviews presented by Akhter et al. [87] and Ahmad et al. [88] elaborated the properties and applications of metallic NPs in oncology.

**Magnetic Nanoparticles**

The basis behind the formulation of magnetic nanoparticles (NPs) is the fact that the drug is either encapsulated into magnetic micro-/nanosphere or conjugated onto them, or implanted as a magnetically-active disc. The drug release to the blood stream can be controlled utilizing a powerful magnetic field in the target tumor area. Magnetic materials receive their magnetic responsiveness to a magnetic field from materials like magnetite, iron, nickel, cobalt, neodymium-iron-boron, and samarium-cobalt. Ferrofluids are liquids which become strongly magnetized in the presence of a magnetic field. They are colloidal liquids of nanoscale ferromagnetic particles suspended in a carrier fluid (organic/aqueous). Iron oxide NPs is the most commonly used due to its biodegradable nature, biocompatibility, super-paramagnetic effects and ability to serve as a contrast agent in MRI. Once compartmentalized within the lysosomes of RES cells, they disintegrate to ferritin and/or haemosiderin (antiferromagnetic forms of iron) [89]. A water-dispersible oleic acid (OA)-pluronic-coated iron oxide magnetic NP formulation was developed by Jain et al. and subsequently loaded with high doses of water-insoluble anticancer agents. This formulation was used as a universal drug carrier system for systemic administration of water-insoluble drugs, and simultaneously permitting magnetic targeting and/or imaging [90]. Yallapu and co-workers have developed a multilayer approach for water-dispersible superparamagnetic iron oxide nanoparticles (SPIONS) for hyperthermia, MRI and drug delivery applications. Iron salts, in the presence of ammonia, were precipitated to obtain iron oxide core NPs and coated with β-cyclodextrin and pluronic polymer. Consequently, the formulation was highly water dispersible, enabling entrapment of anticancer drugs in β-cyclodextrin and pluronic polymer coatings for sustained drug release and showed superior hyperthermia effects with time under alternating magnetic fields as compared to pure magnetic nanoparticles and β-cyclodextrin coated NPs [91]. In yet another study conducted by Maeng et al., multifunctional doxorubicin-loaded SPIONS for chemotherapy and MRI in liver cancer were developed. Polymeric NPs composed of poly(ethylene oxide)-trimellitic anhydride chloride-folate, doxorubicin, superparamagnetic iron oxide and folate, and exhibited superior anticancer activity by targeting folate receptor (FR)-expressing tumors, thus enhancing the bioavailability and efficacy of the drug [92].

**Dendrimers**

Dendrimers form an important connecting link between molecular chemistry and polymer science [92]. They are highly branched, globular, monodisperse, nanoscaled, and uniformly distributed three-dimensional polymeric macromolecules consisting of three distinct domains: (a) a central core containing a single atom/atomic group with at least two identical chemical functions, (b) branches emanating from the core, comprising of repeating units organized in geometric progression resulting in a series of
Radially concentric layers called “generations”, and (c) exteriorly located terminal functional groups. Since they possess both hydrophobic and hydrophilic areas, drug molecules can be loaded depending upon their solubility characteristics. They serve as an ideal targeting and imaging agents due to their highly asymmetric shape, multiple branching and multivalent tree-like structures. Among different dendritic polymers, polyamidoamine (PAMAM) dendrimers are the most widely used for chemotherapeutics. In the recent past, multifunctional PAMAM dendrimers with an imaging agent (fluorescein isothiocyanate) conjugated with biotin and folic acid as targeting ligands, and taxol as therapeutic agent have been elucidated for parallel use in diagnosis and therapy. Conjugation of partially-acetylated PAMAM dendrimers with imaging modalities might be used for cancer therapy and diagnosis. Enhanced drug loading and extended delivery of 5-fluorouracil was observed with pegylated PAMAM dendrimers, with few blood dyscrasias due to restricted uptake of drug through RES [93]. Gardikis et al. developed a liposomal-locked indendrimer (LLD) formed by dipalmitoylphosphatidylcholine-dipalmitoylphosphatidylglycerol (DPPC-DPPG) lipids and poly(amide amine) incorporating the drug doxorubicin. The study showed that phase separation between DPPC-DPPG lipids and dendrimer promote the stability of liposomal membrane and co-operativity of the relevant gel to liquid crystal transitions, which is augmented in the presence of dendrimers and drug [94].

Carbon Nanotubes (CNTs)

Over the last several years, CNTs have been extensively explored in almost every single cancer treatment modality, including drug delivery, lymphatic targeted chemotherapy, thermal therapy, photodynamic therapy and gene therapy. CNTs were first discovered by Iijima in 1991 [95]. They belong to the family of fullerenes, i.e. the third major allotropic form of carbon after graphite and diamond [96]. They are structurally thin sheets of benzene ring carbons rolled up uniformly to form a smooth, seamless rod-like tubular structure. CNTs can be single-walled (SWNT), which essentially consist of a layer of cylinder graphene, or multi-walled (MWNT) containing several concentric graphene sheets [96]. Some important techniques to produce CNTs are laser ablation, electric arc discharge, and thermal or plasma-enhanced chemical vapor deposition (CVD) [97, 98]. Liu et al. synthesized a series of amphiphilic polymers by anchoring PEG of different lengths at various densities on poly(maleic anhydride-alt-1-octadiene) [PMHC18]. PEG-PMHC18-coated SWNTs in mice after i.v. injection showed ultra-long elimination half-life and significantly higher tumor uptake [98]. Chen et al. demonstrated that a higher epirubicin loading can be achieved by utilizing carboxylated CNTs (cMWNT) [99]. The loading process was mainly attributed to π-π stacking between epirubicin and graphene surface of CNTs. Carboxylic acid group also facilitated functionalization with the targeting group that PEGylated MWNTs penetrated mammalian cells without damaging the plasma membrane and selectively accumulated in MDR-cancer cells as efficiently as in sensitive cancer cells [99]. These intracellular translocations of PEGylated MWNTs were visualized in MDR-Hep G2-DR and sensitive Hep G2 cells, as observed by fluorescence and transmission electron microscopy. This study, thus gives the strong evidence regarding development of PEGylated MWNTs as an efficient drug carriers to conjugate drugs for overcoming MDR in cancer chemotherapy [99].

Quantum Dots (QDs)

Semiconductor QDs are nanoparticles (2-100 nm) that are considered as the potential candidate in traceable drug delivery because of their size-tunable absorption bands and emission color [100]. Additionally, photoluminescence of QDs is outstandingly bright and stable, making them suitable alternatives for biomedical imaging and therapeutic interventions [101]. Semiconductors from groups II-IV or III-V of the periodic table are employed, for example, indium arsenide, cadmium telluride, and cadmium selenide. Various advantages of QDs over ionizing radiations, chemotherapy and radiation therapy are: (a) high quantum yield; (b) resistance to chemical modification; (c) intrinsic fluorescence emission spectra; and (d) adjustable optical properties by regulation of size and composition. Photosensitizing QDs produce radicals upon absorption of visible light. This approach is limited for superficial tumors. Another major drawback to the use of QDs is the poor availability of photostable NIR fluorophores and photosensitizing drugs [100]. A QD-aptamer-doxorubicin conjugate functionalizing the surface of fluorescent QDs with A-10 RNA aptamer was developed by Bagalkot et al. These QDs were able to differentially uptake and image prostate cancer expressing prostate-specific membrane antigen [102]. Gao et al. developed multifunctional QD probes for simultaneous tumor targeting and imaging. Early diagnosis and individual tailorized treatment of oral cancers could be achieved by developing NIR QD conjugated with cell penetrating peptide and labelling oral squamous carcinoma cells with QD conjugates [103]. Li et al. utilized PAMAM dendrimers to modify QDs, thereby enhancing their water solubility. These dendrimer-modified QDs were conjugated with DNA aptamers specifically targeted to U-251 human glioblastoma [104]. Chen et al. investigated the potential of QDs for quantitative estimation of HER2 as a micropathologic indicator of tumor size. Usage of QDs has led to a better revelation of breast cancer heterogeneity, which can, in turn be useful in formulating a more personalized and targeted therapy for breast cancer [105].

RECENT ADVANCEMENTS AND FUTURE TRENDS

Attenuation of MDR (Multi-Drug Resistance)

Resistance against drugs remains the major challenge in oncology. Drug resistance can be of non-cellular and cellular origin. Poorly vascularized tumor cells/tissues or physiological barriers are considered as the cause of non-cellular drug resistance that significantly diminishes drug access to the cancerous sites. Cellular drug resistance can be due to over-expression of the drug transport pump (for example P-gp efflux transporter), drug-resistance proteins, enhancement of the DNA repair capacity and reduction of the apoptosis regulation [106]. Amongst the aforementioned mechanisms, the role of P-gp is most extensively studied. P-gp is a 170-kDa transmembrane glycoprotein that functions
as an efflux system to remove xenobiotics from inner cell environment. Various P-gp inhibitors have been discovered to overcome drug resistance (e.g. cyclosporin A, ramipril and cuminaldehyde) and among them, some of the P-gp inhibitors have shown the restoration of cancerous cell sensitivity to cytotoxic agents [106].

Alternative strategies against MDR are the development of the novel concept of nanotechnology-based drug delivery systems, which selectively target the tumor-site (cellular or tissue targeting) through selective endocytosis or passive diffusion leading to by-pass the P-gp efflux [6], for example, doxorubicin-loaded poly-(alkyl cyanoacrylate) NPs [107], metallic doxorubicin-loaded particles, PACA NPs [108] and nanoparticulate conjugate of 20 (S)-camptothecin [109]. Schluep et al. have shown the ability of such particles to overcome drug resistance [109]. In clinical studies, liposome-loaded doxorubicin was able to overcome drug resistance in AIDS-related Kaposi’s sarcoma [110]. Furthermore, NPs such as polymeric micelles and polymeric conjugates as a drug delivery carrier showed promising results in patient who had previously failed to chemotherapy [110-112]. Ligand-based strategies, particularly receptor targeting, have also been applied to overcome MDR since they are able to interact with the receptor and internalized through receptor-mediated endocytosis. For example folate-targeted DOX-nanoparticles and transferring-conjugated paclitaxel NPs exhibited greater cytotoxicity than their free counterpart in MDR model [10].

Nanotechnology in Cancer Theranostics

Diagnosis and therapy combined in one system is the current advanced biomedical tool referred to as theranostics [87]. The primary goal of theranostics is to selectively target the disease confined to tissue/cell in order to increase diagnostic and therapeutic selectivity that makes the treatment shorter, safer, effective and inexpensive. Biocompatible/biodegradable nanocarriers are currently under development in oncological theranostics that would enable precise diagnosis and therapy. Lukianova-Hleb et al. have studied the light-based generation and detection of plasmonic gold NPs in living cells, with focus on tailoring the plasmonic NP properties in one cell and evaluating the multifunctionality of the nanoparticle [113]. Recently, numerous published reports discussed the design, physiochemical characteristics and applications of magnetic nanoparticles. These NPs can act both an imaging agent (diagnosis) and a drug carrier (therapy) [114]. Shim et al. coated RNA-encapsulated polyplexes, covalently bonded with gold nanoparticle through an acid labile linkage to explore theranostic use (optical imaging and gene silencing) [115]. The different possible characteristics like imaging property, drug carrying and targeting capacity with long circulating behavior (shown with PEG) are given in Fig. (2).

The multifunctionality associated with NPs includes imaging (single or dual modality), therapy (single or multiple drugs) and targeting (uni- or multi-liganded). Although there are certain unanswered questions and challenges remaining for the establishment of NPs in clinical applications, extensive clinical data are needed, with progressive and clinical-centric efforts to overcome these queries and obstructions that will certainly lead to rational design of nanoparticles, possessing improved selectivity and efficacy. Current knowledge regarding the safety of NPs is in its infancy. The pharmacokinetic parameters and their outcome need a comprehensive exploration and generation of databases for health-risk particularly associated with vital organs need to be created. In the next few years, many applications of nanotechnology will be put to clinical practice. Table 3 summarizes the selected examples of nanomedicines which are approved by the FDA or are in some phase of clinical trial for cancer therapy.

TOXICITY ISSUES

Apart from immense importance of nanomaterials in cancer imaging and therapy, its toxic effects are also a major concern especially after chronic administration. Different polymers have different toxicities, like complement activation, carcinogenicity, teratogenicity, and immunogenicity [116, 117]. Thus choosing safe polymers for the design of NPs is itself a major hurdle. Careful evaluation of
the potential toxicity of residual solvents, polymers and the developed particles is critically important. Presently, nontoxic and biodegradable ingredients are used to design NPs due to which carrier-associated toxicities tend to be mild. However, NPs accumulates in the liver, spleen, and bone marrow leading to increased toxicities to these organs. Many studies have identified liver as the primary organ responsible for reticuloendothelial capture of NPs, often due to phagocytosis by Kupffer cells. Hepatotoxicity has been observed in mice treated orally with nano-zinc particles [118]. Similarly, i.v. administration of cationic PAMAM dendrimers to mice has been observed to cause liver injury [119]. There are also safety concerns with particular NPs that are capable of crossing the BBB. For example clinical trial of an HPMA-conjugated paclitaxel had been terminated due to neurotoxicity [120]. Among the nanomedicines, metallic nano-carriers have been extensively evaluated for their toxicity and interaction with biological systems [121]. Depending on the nature and type of metallic nanoparticles (MNPs), different grades of toxicity have been reported at

### Table 3. Nanomedicine as cancer therapeutics/imaging agents under different stages of development.

<table>
<thead>
<tr>
<th>NANOMEDICINES</th>
<th>DRUG</th>
<th>APPLICATION</th>
<th>STATUS</th>
<th>REFERENCE/COMPANY</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LIPOSOMES</strong></td>
<td>Paclitaxel</td>
<td>Malignant pleural effusions</td>
<td>Phase I</td>
<td>NeoPharm, USA</td>
</tr>
<tr>
<td></td>
<td>CPX-1 (Liposomal irinotecan)</td>
<td>Colorectal cancer</td>
<td>Phase II</td>
<td>Celator Pharmaceuticals</td>
</tr>
<tr>
<td></td>
<td>INGN-401</td>
<td>Metastatic lung cancer</td>
<td>Phase I</td>
<td>Introgen Therapeutics Inc, USA</td>
</tr>
<tr>
<td></td>
<td>SPI-77 (stealth liposome cisplatin)</td>
<td>Lung cancer</td>
<td>Phase III</td>
<td>ALZA pharmaceuticals, USA</td>
</tr>
<tr>
<td></td>
<td>Cytosine arabinoside</td>
<td>Neoplastic meningitis</td>
<td>Phase IV</td>
<td>ALZA pharmaceuticals, USA</td>
</tr>
<tr>
<td></td>
<td>Daunorubicin citrate</td>
<td>Kaposi’s sarcoma</td>
<td>FDA approved</td>
<td>Gilead Sciences, Inc,USA</td>
</tr>
<tr>
<td></td>
<td>Doxorubicin</td>
<td>Ovarian cancer</td>
<td>FDA approved</td>
<td>ALZA pharmaceuticals, USA</td>
</tr>
<tr>
<td></td>
<td>PEGylated liposome/doxorubicin hydrochloride</td>
<td>Ovarian cancer</td>
<td></td>
<td>Ortho Biotech</td>
</tr>
<tr>
<td></td>
<td>Gentuzumab ozogamicin</td>
<td>Acute myeloid leukemia</td>
<td>FDA approved</td>
<td>Wyeth-Ayerst, USA</td>
</tr>
<tr>
<td></td>
<td>Vincristine sulfate</td>
<td>Acute lymphoblastic leukemia</td>
<td>FDA approved</td>
<td>Hana Biosciences, Inc, USA</td>
</tr>
<tr>
<td></td>
<td>OSI-211 (Liposomal lurtotecan)</td>
<td>Various cancer</td>
<td>Phase II</td>
<td>OSI Pharmaceuticals</td>
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<tr>
<td><strong>POLYMERIC MICELLES</strong></td>
<td>MRX-952</td>
<td>Anticancer</td>
<td>Preclinical</td>
<td>ImaRx Therapeutics, USA</td>
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<td></td>
<td>Carmustine</td>
<td>Glioblastoma multiforme</td>
<td>FDA approved</td>
<td>Guilford Pharm.Inc. USA</td>
</tr>
<tr>
<td><strong>SOLID-LIPID NANOPARTICLES</strong></td>
<td>Paclitaxel-taxol</td>
<td>Mammary cancer (metastatic)</td>
<td>FDA approved</td>
<td>American Pharm.Partner/Amer. BioScience, USA</td>
</tr>
<tr>
<td><strong>POLYMERIC NANOPARTICLES</strong></td>
<td>Copolymer of N-(2-hydroxypropyl) metacrylamide/camptothecin</td>
<td>Various cancer</td>
<td>Phase I</td>
<td>Pharmacia, USA</td>
</tr>
<tr>
<td></td>
<td>NK-105 (PEG-polyaspartate) nanoparticle of cisplatin</td>
<td>Various cancer</td>
<td>Phase II</td>
<td>Nippon Kayaku Co. Ltd.</td>
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<tr>
<td><strong>IRON OXIDE NANOPARTICLES</strong></td>
<td>Combidex</td>
<td>Tumor imaging</td>
<td>Phase III</td>
<td>Advanced Magnetics, USA</td>
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<td></td>
<td>Resovist® (Iron oxide nanoparticles coated with carboxydextran)</td>
<td>Tumor imaging</td>
<td>In European market</td>
<td>Bayer Schering Pharma AG</td>
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<tr>
<td></td>
<td>Feridex® (Iron oxide nanoparticles coated with dextran)</td>
<td>Tumor imaging</td>
<td>FDA approved</td>
<td>Berlex Laboratories</td>
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<td><strong>ALBUMIN NANOPARTICLES</strong></td>
<td>Albumin bond nanoparticle (Abraxane)</td>
<td>Mammary cancer (metastatic)</td>
<td>FDA approved</td>
<td>American Pharm.Partner/Amer. BioScience, USA</td>
</tr>
<tr>
<td><strong>CYCLODEXTRIN NANOPARTICLES</strong></td>
<td>Cycloser-t-camptothecin</td>
<td>Metastatic solid tumors</td>
<td>IND filed</td>
<td>Insert Therapeutics, USA</td>
</tr>
<tr>
<td><strong>GOLD NANOPARTICLES</strong></td>
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<td>Diagnostics</td>
<td>Market</td>
<td>Nanosphere, USA</td>
</tr>
<tr>
<td></td>
<td>Aurimune (Colloidal gold/TNF)</td>
<td>Solid tumors</td>
<td>Phase II</td>
<td>CytoImmune Science</td>
</tr>
<tr>
<td><strong>SILICA NANOPARTICLES</strong></td>
<td>AuroLase (Gold coated silica nanoparticles)</td>
<td>Head and neck cancer</td>
<td>Phase I</td>
<td>Nanospectra bioscience</td>
</tr>
</tbody>
</table>
organ, tissue, cellular and sub-cellular level due to their physiochemical properties (viz. size, shape, electrical charge on the surface, chemical composition, surface structure (surface reactivity, surface group, inorganic or organic coatings), solubility and aggregation behavior) [122-124]. The mechanisms of toxicity induced by MNPs are a combination of different events such as direct destruction of cellular component like DNA, RNA and cellular proteins due to free radical generation. Moreover, increased oxidative stress plays a key role as well [125]. Recently, it was found that ‘naked’ CdTe quantum dots cause damage to lipid membranes of mitochondria and nuclear structures [125]. Destruction of plasma membrane by metallic NPs was supported by the work of Sayes et al. on Cobalt (60Co) NPs [126]. Metallic NPs influence the reproductive organs as it was found that particles may reduce spermatogonial cell proliferation [127-130]. Free radical generation and induction of inflammatory mediators is considered pivotal for such effects [131]. Apart from the male reproductive organ, Browning et al. reported the effect of metallic NPs over female reproductive system in animal experiments [132]. They confirmed that gold NPs can diffuse into the embryo and lead to teratogenic deformities [133]. Increased surface activity due to reduction in particle size (increase in surface area:volume ratio) influences the interaction of metallic NPs to the biological system [87]. The correlation between the size and toxicity is corroborated by the report of De and co-workers that illustrated that colloidal gold NPs with small size (10-50 nm) cause more toxicity in comparison to the larger particles (100-200 nm) [134]. Moreover, Chen et al. studied the oral toxicity of copper NPs and found that their LD50 increased sharply with the decrease in particle size [135]. In another report, it was demonstrated that gold NPs with the size ranging from 2.8 to 38 nm were more toxic and induced immunological reactions [136]. Size-dependent adverse effects of silver NPs were found in in-vitro studies on hepatocytes indicating decreased mitochondrial function, lactate dehydrogenase (LDG) leakage and abnormal cell morphologies [137]. Influence of surface charge on the metallic NPs has also been addressed in the literature. Recently, dose-dependent effect on viability and capacity of nerve cells was seen in case of anionic MNPs [138]. Although in vivo findings on MNPs toxicity have also been reported, but majority of the data on the toxicity of MNPs have been the outcome of in vitro testing. It is promising that various efforts are being undertaken for reducing the uptake of NPs by macrophages (MPs) and to increase their accumulation at the active site through surface modification and/or incorporation of targeting ligands to the polymers/NPs [139-141]. On the basis of reports, the toxic

Fig. (3). Simplified pictorial diagram representing the possible toxic effects of nanomedicines on different body organs.
effects of nanomedicines on different body organs are exemplified in Fig. (3). In nanomedicine formulation development, PEGylation is the most common technique used to make long circulating nano-carriers. Moreover, plenty of published papers on this subject indicate that such hydrophilic polymer coating reduces the nanoparticles-associated toxicity. Regardless of the prevalent use of PEGylation for prolonging the circulation of nanomedicines, its mechanism for such action and interaction with biological system in the systemic circulation is not entirely clear. In fact, PEG has been shown to form a brush border over the nanomedicines’ surface that leads to non-specific impermeable hindrances and sterically prevents access of tissue proteins [142]. Chain length, density, conformational freedom and flexibility of PEG are critical in making the nanomedicine-surface free from cellular interaction [143]. Although, such kind of coating provides non-specific shielding, but it was found that in case of PEGylated hexadecyl cyanoacrylate NPs, PEG was able to reduce the adsorption of ApoC and immunoglobulins, but not the other plasma proteins [144]. Similarly, with uncharged liposomes (made up of phosphatidylcholine and cholesterol) and negatively charged liposomes (made up of phosphatidic acid), coating with PEG-DSPE 2000 does not provide shielding from plasma protein adsorption [145]. Recently, several reports have been published that indicate the rapid clearance of PEGylated liposomes [146]. This effect is attributed to the formation of PEG-specific antibodies that prompt complement absorption [145]. Therefore, it should be noted that expression of specific antibodies to the particles or coating material may affect the pharmacokinetics, efficacy and safety of nanomedicines.

Toxicity Assessment

In Vitro Cells and Gene-Based Toxicity

To generate information on toxicity, studies at different levels (in vitro, in vivo and biochemical) must be carried out. Although in vitro studies may overcome the complexity of animal-based experiments, they may oversimplify processes and lack in vitro-in vivo correlation. However, they allow for a level of control well-suited for elucidating toxic effects and their mechanism at molecular level, which is extremely difficult to interpret in in-vivo studies. A gene-based approach (DNA strand breaks, chromosomal damages and gene mutation, etc.) has the advantage of assessing toxicity at low-dose level and may detect carcinogenic effects which may not be monitored via classical cytotoxicity studies. The nature and mechanism of interaction between NPs, and molecular and cellular targets can be elucidated by TEM, SEM or X-ray-based microscopy.

Oxidative Stress and Reactive Oxygen Species (ROS) Assessment

Reduction of size to nano-scale leads to the conversion of normal particles to highly-active charged particles, which on interaction with cellular components and biological processes generate ROS, and produce oxidant injury. This paradigm may be used to evaluate the toxic potential of NPs [87]. Oxidative stress occurs when ROS production exceeds the defense capacity of the system and in such circumstances leads to damage of biomolecules like lipids, DNA, RNA, and proteins, resulting in excessive cell proliferation, apoptosis, and mutations. Metallic and inorganic NPs are known to induce oxidative stress through ROS generation during redox reaction by interruption of the electronic flux, perturbation of the permeability transition, and diminution of protective cellular component like glutathione [146-148]. ROS production can be monitored through ESR or with the use of fluorescent spectroscopy with or without quenchers such as furfuryl alcohol and superoxide dismutase. Furthermore, within a living cell this activity can be monitored through fluorescent microscopy or confocal microscopy. In addition, change in intracellular free calcium, mitochondrial function and structural integrity also serves as a tool to study ROS production and their associated oxidative stress.

REGULATORY ASPECTS

The positive impact of nanomedicines on the quality of life in life-threatening diseases like cancer is evident in clinical trials. However, applying these initial outcomes to clinical practice still needs to be rationalized and carefully controlled in terms of risk-to-benefit ratio, and is carefully overseen by regulatory authorities such as FDA, MHRA and EMEA. For any such products, the agencies have to set protocols that basically work for the assessment of three parameters: quality, safety and efficacy. The first generation of nanomedicines is already in clinical practice and some of it is now in the phase of generic development. Both generic manufacturers and regulatory agencies are struggling to finalize the studies that are required to demonstrate the bio-equivalency of nanometric generic medicines compared to innovator products. This has been clearly demonstrated with several unproductive efforts in the generic development of nab-paclitaxel formulation. The claimed bio-similar of nab-paclitaxel, when evaluated, did not fit to reproduce particle size, stability, potency and other physicochemical attributes of nab-paclitaxel [149]. Moreover, unlike nab-paclitaxel, the reconstituted nanomedicine exhibited poor stability when evaluated for accelerated stability testing i.e. it formed aggregates within 24 h of study period. Pharmaceutical or chemical equivalence and/or bioequivalence may not sufficiently indicate the function of nanomedicines at the site of action, as assumed for standard conventional preparations. In addition, several liposomal formulations containing amphotericin B and doxorubicin have recently gone off-patent. The lack of critical information regarding composition, dimensional configuration of components and critical parameters that are essential for optimal performance of nanomedicines, raise concern about “generics” that may be approved based on conventional drug product standard and the guidelines but without having the same quality of the innovator. This exemplifies that how generic nanomedicine manufacturers and regulatory authorities are going to face challenges in the development and approval of such products. So, it is crucial that a comprehensive physicochemical based understanding of nanomedicines and recognition of critical parameters that affect their functioning be conducted early in developmental that will help regulatory bodies and industry in framing quality standard of nanomedicines products. Indeed, FDA has recently begun to consider relevant approval standards for generic copies of nanomedicines. A recently issued guideline in case of
doxorubicin-loaded liposomal formulation (Doxil®) is an example for the generic development. Current scenarios illustrate that nanomedicines have defined their physico-chemical characteristics but lack suitable pharmacokinetic models for such medicines. Considerations such as pathophysiology, target tissue, biodistribution and elimination, impact of their size and surface characteristics on organ, tissue and cellular localization, and better understanding of pharmacokinetics and pharmacodynamics correlations need to be addressed in experimental models. Moreover, quantitative techniques need to use for biodistribution studies of polymers and metals. In the context of polymeric and metallic nanomedicine therapeutics, carrier systems are developed with newer complex architectures (for example, dendrimers, quantum dots, SPIONS and multifunctional gold nanomaterials, etc.) and such carrier systems may be given by different route, viz pulmonary, i.v. and organ-directed injections. With the increased complexity of the architecture, induction of multifunctionality in a single carrier system frequently falls into a gap between medicines and medical devices regulation. It is to be noted that biological and medical devices assessment guidelines are based on general and non-specific standards. Moreover, a regulatory guideline addressing new nanometric devices and different category of medicines harmonized for the global regulatory will be highly productive.

CONCLUSION

In the current multidisciplinary arena, nanotechnology is in a distinctive status to transform cancer chemotherapy and diagnosis to produce a new generation of cancer therapeutics/theranostics (categorized as nanomedicines) with high sensitivity and precision for cancerous cells leading to reduction of the conventional therapeutics-associated predictable toxicity. There are an increased number of FDA-approved cancer nanomedicines, their socio-environmental impact need to be addressed. While establishing the nanomedicines in clinical oncology practice, some key points need to be considered; toxicological data review of nanomedicines prior to use, in vivo-in vitro toxicity testing to understand the toxicity of new nanomedicines and its environmental impact.

ABBREVIATIONS

AML = Acute Myeloid Leukemia, Acute Myelogenous Leukemia
BBB = Blood-Brain Barrier
CNTs = Carbon nanotubes
CVD = Chemical Vapor Deposition
DOX = Doxorubicin
DPPC-DPPG = Dipalmitoylphosphatidylcholine - Dipalmitoylphosphatidylglycerol
EPR = Enhanced Permeability and Retention
HER2 = Human Epidermal growth factor Receptor 2
IGF-1R = Insulin-like Growth Factor-1 Receptor
MDR = Multi Drug Resistance
MM = Multiple Myeloma
MWNT = Multi-walled Carbon Nanotubes
NSVs = Nonionic Surfactant Based Vesicles
PACA = poly(alkyl cyanoacrylate)
PAMAM = Polyamidoamine
PGP = P-glycoprotein
PLGA = poly(lactic-co-glycolic acid)
PSMA = Prostate-Specific Membrane Antigen
RES = Reticuloendothelial System
ROS = Reactive Oxygen Species
SWNT = Single-walled Carbon Nanotubes
TMNPs = Theranostic Metallic Nanoparticle
TNF = Tumor Necrosis Factor
VEGF = Vascular Endothelial Growth Factor
VIP = Vasactive Intestinal Peptide
WGA = Wheat-Germ Agglutinin
WIT-NP = Wheat-Germ Agglutinin Nanoparticles

CONFLICT OF INTEREST

The author(s) confirm that this article content has no conflicts of interest.

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