Potential applications of nanotechnology for the diagnosis and treatment of pancreatic cancer

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INTRODUCTION

Pancreatic cancer is one of the most lethal cancers worldwide with a 5-year relative survival rate of <6% (Jemal et al., 2011; Siegel et al., 2013). Despite aggressive combinations of therapies ranging from surgery, chemotherapy, and radiotherapy, patients diagnosed with this malignancy have extremely high mortality rates and poor prognosis (Vincent et al., 2011). These dismal outcomes can in part be attributed to a lack of early diagnosis and inability to detect pre-cancerous pancreatic intraductal neoplastic (PanIN) lesions, which often give rise to invasive pancreatic tumors (Misek et al., 2007). Currently no clinically reliable serum biomarkers for early detection and diagnosis at an early stage of pancreatic cancer are available (Misek et al., 2007). A majority of pancreatic cancers are diagnosed and staged by computed tomography (CT) imaging or magnetic resonance imaging (MRI) with a tumor detection limit of 5–8 mm when the earliest precursor lesions are in the microscopic range (Holzapfel et al., 2011; Canto et al., 2012). Consequently, >80% of patients have locally advanced or distant metastases upon diagnosis, with chemotherapy or palliative chemotherapy being the current standard for systemic treatment (Vincent et al., 2011). Notably, only 20% of patients at the time of diagnosis are suitable for potentially curative pancreatectomy and despite combination with adjuvant chemotherapy, a majority of patients still develop local recurrences and systemic metastases that results in mortality (Hidalgo, 2010). Another contributing factor to the poor outcome of pancreatic cancer is that the current standard first-line therapy of the chemotherapeutic agent gemcitabine shows only limited efficacy by extending the overall survival of patients by only 6–12 weeks (Hidalgo, 2010). This is partially due to poor understanding of the biology of the individual patients’ tumor, chemotherapy resistance and the tumor microenvironment.

Nanotechnology is defined as the manipulation of organic or inorganic materials to form structures on the scale of nanometers. Recently, advances in nanotechnology have provided great opportunities for strategies in advancing cancer diagnostics, imaging, and therapeutic drug delivery (Schroeder et al., 2011;...
Melancon et al., 2012; Prabhu and Patravale, 2012). Nanoparticles have the potential to not only increase the efficacy per dose of a therapeutic or imaging contrast formulation by increasing its bioavailability, but can also be modified for targeted selectivity toward tumor cells to increase image resolution and/or reduce off-target toxicities associated with current chemotherapy (Ferrari, 2005). Furthermore, they show promise in treating metastatic cancers like pancreatic cancer [see detailed review (Schroeder et al., 2011)]. Nevertheless, in order to design effective nanoparticles to deliver either imaging contrast agents or therapeutics (by passive or active targeting) requires a thorough understanding of the physiological barriers specifically associated with the disease and apply nano-engineered components for effective nanoparticle extravasation, accumulation and penetration into the tumor (Jain and Stylianopoulos, 2010). This review highlights promising translational prospects and the challenges of integrating nano-engineering sciences with pancreatic cancer biology to develop nanomaterials that could enhance diagnostics, imaging, and therapeutics against this devastating malignancy (Table 1).

THE PANCREATIC TUMOR MICROENVIRONMENT AND ITS ROLE IN PROMOTING CHEMOTHERAPY RESISTANCE

Desmoplasia and hypovascularity are the pathological hallmarks of pancreatic tumors. The desmoplastic microenvironment (also known as the stroma) can make up >90% of the tumor mass (Li et al., 2010; Neesse et al., 2011). The stroma contains a number of different cell types including, pancreatic stellate cells (PSCs), endothelial cells, immune cells and dense extracellular matrix (ECM) (Li et al., 2010; Neesse et al., 2011). Indeed, it is well established that PSCs are the principle cell type responsible for the production of stromal fibrosis (Apte et al., 2004; Bachem et al., 2005). Importantly, the extensive fibrotic characteristic of pancreatic cancer results in reduced intratumoral vascular density which give rise to compromised dysfunctional vessels that cause a decrease in blood flow; inadequate venous and lymphatic drainage also further increases the interstitial fluid pressure within pancreatic tumors (Koong et al., 2000; Erkan et al., 2007; Komar et al., 2009; Olive et al., 2009). As a consequence the stroma has been shown to play a major role in poor chemotherapy drug delivery, penetration, and rapid metabolic inactivation of therapeutic agents which contribute to an unusually poor response to treatment (Olive et al., 2009; Provenzano et al., 2012). Furthermore, the bi-directional interaction that occurs between PSCs and the tumor cells further potentiates tumor progression, chemoresistance, invasion, and metastases (Apte et al., 2004; Bachem et al., 2005; Vonlaufen et al., 2008; Olive et al., 2009; Xu et al., 2010; Phillips, 2012). However, the stroma could also be the achilles’ heel for targeted diagnostic imaging and therapeutic drug therapy to ablate the microenvironment that supports tumor growth and metastases (Phillips, 2012; Heinemann et al., 2013). In addition, given the tumor mass comprises mainly of the fibrotic stroma, the ECM components or PSCs residing within the stroma could be novel targets for early diagnosis, imaging, and targeted therapy. Examples of how nanotechnology may be applied to advance pancreatic cancer diagnosis and treatment are described below.

NANO-DIAGNOSTICS FOR THE EARLY DETECTION OF PANCREATIC CANCER

The vast majority of long-term survivors of pancreatic cancer (> 5 years) have resectable disease upon diagnosis and are suitable for curative surgery, suggesting early detection and intervention may increase the overall survival of patients (Slavin et al., 1999). However, currently there are no reliable serum biomarkers with the sensitivity and specificity to accurately detect early pre-cancerous lesions (Goggins, 2005). This is largely due to the lack of pancreatic cancer biomarkers able to distinguish between a benign diseased pancreas, such is the case for chronic pancreatitis and cancer, as both are hard to distinguish under current imaging modalities (Erkan et al., 2012a,b). Moreover, the heterogeneous nature of pancreatic cancer and the complex stromal microenvironment also present a challenge for potential biomarkers. Hence, early diagnosis of pancreatic tumors may require the simultaneous identification of a panel of biomarkers to have greater accuracy compared to single biomarkers.

Nano-diagnostic platforms have the potential to revolutionize the cancer diagnostics field by developing faster, more accurate, cost-effective, and reliable biomarker detection systems with lower detection limits (Cao, 2008; Chikkaveeraiah et al., 2012; Malhotra et al., 2012). Various electrochemical immunosensors have also integrated nanomaterials such as magnetic particles, gold nanoparticles, quantum dots, and carbon nanotubes to increase their sensitivity for electrochemical detection of tumor biomarkers (Cao, 2008; Chikkaveeraiah et al., 2012; Malhotra et al., 2012). For instance, microfluidic biochips integrated with highly luminescent quantum dots have shown promise as a versatile multicolor and multiplexed bioassay (Hu et al., 2010). This was evidenced by their ability to rapidly detect two cancer
biomarkers, carcinoma embryonic antigen, and α-fetoprotein, with high selectivity and femtomolar sensitivity in human serum (Hu et al., 2010). Recently, Chang et al. (2011) managed to design passivated (technique used to prevent or retard non-specific background signal) nanowire biosensors to allow rapid and specific detection of the ovarian cancer biomarkers, cancer antigen-125 and insulin-like growth factor-II directly from human whole blood collected simply from a finger prick, with much lower limits of detection compared to the clinically relevant levels for diagnosis (Chang et al., 2011). Others like, the NanoMonitor incorporated nanoporous alumina membranes onto micro-fabricated silicon platforms allowing rapid label-free analysis of glycans from pancreatic cancer cell lysates with high sensitivity and selectivity (Nagaraj et al., 2010). Biosensors like these could be useful for pancreatic cancer diagnosis in the clinic by identifying other transmembrane glycoproteins such as mucins (MUC) which are expressed on the surface of pancreatic cancer cells and are important regulators of tumor growth and metastases (Kaur et al., 2013). Indeed, studies have demonstrated several members of the MUC protein family to be overexpressed and/or aberrantly glycosylated during the progression of early PanIN lesions to pancreatic tumors in a mouse transgenic model (Kras<sup>G12D</sup>;Pdx1-Cre), which mimics the human setting for the initiation and progression of pancreatic cancer (Rachagani et al., 2012; Remmers et al., 2013). Nano-diagnostics could also be explored in the case of pancreatic cancer-induced paraneoplastic diabetes to aid early diagnosis (Sah et al., 2013). It has been suggested that up to 85% of pancreatic cancer patients have diabetes or hyperglycemia which can manifest as early as 2–3 years before the development of pancreatic cancer (Sah et al., 2013). Hence, nano-diagnostics which employ the use of a combination of early cancer biomarkers along with indicators of β cell dysfunction (resulting in worsening type 2 diabetes and increased adrenomedullin levels) could have potential diagnostic utility, to be explored in a large patient cohort (Hart et al., 2011; Aggarwal et al., 2012; Sah et al., 2013). In summary, nano-diagnostics are powerful tools for cancer detection, prevention, and diagnosis. Nano-diagnostic platforms may provide additional sensitivity with relatively small sample volumes for biomarker detection. Furthermore, the increase in sensitivity, specificity, and multiplex properties for the detection of cancer biomarkers using nano-diagnostics could also be used as a monitoring platform to detect early recurrences or to follow the response of patients to treatment that would otherwise be undetectable with conventional assays.

**NANOMATERIALS FOR ADVANCING PANCREATIC CANCER IMAGING**

Currently, the most accurate initial screening modalities for the detection of pancreatic cancer are endoscopic ultrasonography (EUS) and/or MRI cholangiopancreatography (MRCP) (Hekimoglu et al., 2008; Hyodo et al., 2012). The limitation of current imaging-based screens is that upon identification of potentially benign cysts/lesions, subsequent invasive evaluation by collecting tissue biopsies (for example through fine needle aspiration) is still required to confirm diagnosis (Brand, 2001). Early diagnostics to identify carcinomas in situ like PanIN lesions are often challenging due to the detection limit of current radiological methods and how chronic pancreatitis masses show similar pathological aspects (fibrotic stroma) to pancreatic cancer (Erkan et al., 2012a,b). Only cystic tumors are detectable at a pre-invasive stage with conventional radiological methods due to their large size and high contrast compared to the normal pancreas (Erkan et al., 2012a,b). Furthermore, smaller (<1 cm) metastatic tumor deposits in the liver and/or peritoneal cavity may be overlooked by MRI or CT.

To improve the detection limit of MRI or CT-imaging, an extraneous imaging agent can be administered to the patient to enhance the imaging signals leading to the commercial development of a large range of small organic contrast agents. For example, more than one third of MRI scans are performed in conjunction with a contrast imaging agent. The first generation of organic contrast agents had some limitations (poor specificity and rapid renal clearance) (Roberts, 1997; Kim et al., 2011). To overcome these limitations, contrast agents have been designed in a nanoparticle formulation (Hahn et al., 2011). The incorporation of contrast agents into nanoparticles can offer significant advantages, such as easy functionalization by targeting moieties, higher sensitivity compared to small organic contrast agents and improved biodistribution. For instance, inorganic nanomaterials [including gold nanoparticles, and superparamagnetic iron oxide nanoparticles (SPION)] have been investigated as potential contrast agents for cancer imaging (Godin et al., 2011; Huang et al., 2011). Gold and iron oxide nanoparticles (IONPs) in particular offer several advantages compared to conventional small organic MRI and CT contrast agents (Godin et al., 2011; Huang et al., 2011). For example, IONPs have a lower toxicity profile when compared to gadolinium based contrast agents used in current MRI (Boyer et al., 2010). These nanoparticles have the potential to enhance the contrast between the delayed uptake (hypoperfusion) of the hypovascular tumors when compared to the normal parenchyma during the arterial and venous phase with conventional radiology approaches (Erkan et al., 2012a,b).

Indeed, various nanoparticle systems have been generated with enhanced longitudinal relaxation time (T<sub>1</sub>) and transverse relaxation time (T<sub>2</sub>) for high spatial resolution and simultaneous extraction of physiological, molecular, and anatomical information using MRI. Gadolinium-based contrast nanoparticles have been investigated to enhance T<sub>1</sub>-weighted imaging; while various iron oxides, alloyed and bimetallic ferrite nanoparticles have been shown to be promising T<sub>2</sub> contrast agents (Huang et al., 2011). In a study by Kumagai et al. (2010) the authors designed and synthesized a high-density pegylated-coated iron oxide-gold core shell nanoparticle for MRI imaging. The nanoparticle was approximately 25 nm in diameter. Systemic administration of the nanoparticle to mice bearing a subcutaneous colon tumor or an orthotopic pancreatic tumor resulted in its high accumulation into tumor tissue and low non-specific accumulation in the liver and spleen. In another example, extremely small sized (<4 nm) IONPs have been used as T<sub>1</sub> contrast agents with further improvements in reducing non-specific toxicity and enhanced properties for overcoming the commonly observed “blooming effect” (a phenomenon observed which exaggerates the size of a labeled area as well as blurs the image) of T<sub>2</sub> contrast agents (Kim et al., 2011). In addition, multifunctional nanoparticles
or hybrid systems have also shown great promise. These nanoparticles possess greater signal amplification further improving the diagnostics and imaging sensitivity, while also having the capacity to be used as a therapeutic. For instance, Kirui et al. (2013) reported the use of a gold-iron oxide hybrid nanoparticle to image and treat colorectal tumors in mice. The iron oxide enabled for MR imaging of the tumor, while the gold allowed for photothermal therapy. To improve tumor imaging and targeted therapy the authors conjugated an antibody to the surface of the nanoparticle which recognized the A33 antigen which is expressed on the surface of colorectal tumor cells (Kirui et al., 2013). Importantly, administration of the hybrid nanoparticle to mice with xenografted colorectal tumors showed highly effective tumor imaging and increased tumor cell death when laser irradiation was applied (Kirui et al., 2013). In another example, Kaida et al. (2010) demonstrated the use of a supramolecular nanocarrier system which incorporated a clinically approved gadolinium-based MRI imaging contrast agent and a platinum anti-cancer drug (oxaliplatin). The authors showed successful imaging and combined anti-tumor therapy using the supramolecular nanoparticle in an orthotopic pancreatic cancer mouse model without any significant off-target toxicity (Kaida et al., 2010). Moreover, the amount of gadolinium delivered by the nanoparticle to the tumor was seven times higher when compared to free-gadolinium alone (Kaida et al., 2010). However, it is to be noted, that most nano-formulated contrast agents depend on passive targeting to the tumor site via the enhanced permeability and retention (EPR) effect (Cheng et al., 2013). The EPR effect was first described in 1986 by Matsumura and Maeda (1986) and is a result of rapid and uncontrolled angiogenesis during solid tumor growth which gives rise to leaky and damaged vasculatures within the tumor microenvironment compared to normal tissue (Maeda et al., 2000). This phenomenon allows for vascular extravasation of nanoparticles from a wide size range (100–700 nm) to solid tumors. However, it is to be noted that nanoparticles which enter the solid tumor do not necessarily bind specifically to tumor cells, and have the potential to be taken up by other cell types in the tumor microenvironment (Olive et al., 2009; Erkan et al., 2012a,b).

Nevertheless, despite potential pitfalls with passive tumor targeting, nanotechnology has the ability to advance molecular-targeted imaging in pancreatic cancer owing to the ease of functionalizing nanoparticle surfaces with targeting moieties (antibodies, aptamers, and small molecules) to provide enhanced binding affinity and specificity toward the tumor. This was recently illustrated in a study by Yang et al. (2009) which demonstrated the use of multifunctional nanoparticles to target a cell surface receptor urokinase plasminogen activator receptor (uPAR) which is highly expressed in pancreatic cancer cells and tumor stromal cells. Importantly, the nanoparticles enhanced visualization of pancreatic tumors with a high level of sensitivity in an orthotopic pancreatic cancer mouse model using either non-invasive near-infrared optical imaging or MRI (Yang et al., 2009). In another study, a peptide phage display library was used to screen for small peptides which selectively bound to the surface of pancreatic tumor cells in a genetically-engineered mouse model of pancreatic ductal adenocarcinoma (Kelly et al., 2008). Using this technology the authors identified a small peptide which was able to distinguish the tumor cells from surrounding normal pancreatic ductal cells. Furthermore, proteomic analysis revealed that the peptide bound to plectin-1 (intermediate filament of the cell cytoskeleton). The peptide was then conjugated to the surface of magneto-fluorescent nanoparticles and in conjunction with intravital microscopy and MRI imaging the nanoparticles were able to selectivity detect small pancreatic tumors and pre-cursor lesions with high sensitivity in the mouse model (Kelly et al., 2008).

This study highlights the potential of targeted nanoparticles to allow for the visualization of molecular markers that identify specific stages of pancreatic tumor development. More recently, attempts have been made to identify potential markers for the active targeting of pancreatic stroma. Directing contrast agents to the stromal components of pancreatic cancer may amplify contrast signals at both the tumor and precursor lesion sites. For example, Erkan et al. (2007) showed that peristin a secretory protein that accumulates in fibrotic areas is exclusively produced by activated PSCs and is highly expressed in pancreatic cancer compared to normal pancreatic tissue. Thus, the exclusive expression of peristin in the pancreatic tumor microenvironment may be a potential novel target for molecular imaging of pancreatic cancer (Erkan et al., 2007). This concept was recently illustrated by Eck et al. (2008) which took advantage of the strong light scattering signal from gold nanoparticles. To target tumor stroma the nanoparticles were conjugated with antibodies directed against fibroblast activation protein-α which is produced specifically by activated fibroblasts in tumor stroma (Eck et al., 2008). Taken together, these studies highlight the potential for the combined use of molecular markers which target pancreatic cancer cells and/or the surrounding tumor stroma and nanotechnology to improve the specificity and sensitivity of current pancreatic cancer imaging modalities.

**NANOMEDICINES AS A NOVEL CLASS OF THERAPEUTICS FOR PANCREATIC CANCER**

The design and synthesis of nanoparticles which can encapsulate and deliver a diverse range of therapeutic compounds—ranging from chemotherapy agents to DNA/RNA has received significant attention in cancer research. Nanoparticles in the form of liposomes and/or polymer-derived nanomaterials have been widely used in a number of pre-clinical cancer models. Importantly, these nanoparticles have shown great potential as highly efficient delivery vehicles for chemotherapy drugs or RNA interference (RNAi) inhibitors and are currently being evaluated in human clinical trials (Blanco et al., 2011; Namiki et al., 2011; Schroeder et al., 2011; Singh et al., 2012). A select number of examples for the use of nanoparticles to deliver chemotherapy agents or RNAi inhibitors are described in the following sections.

**NANOPARTICLES AS DELIVERY VEHICLES FOR CHEMOTHERAPY DRUGS**

Many chemotherapeutic agents are associated with debilitating off-target toxicity, poor tumor bioavailability, and unfavorable pharmacokinetics. One strategy to overcome these challenges is the use of nanotechnology as efficient carriers for
chemotherapeutic drugs. The rational design of nanoparticles for chemotherapeutic drug delivery has enabled the improved solubilization of the drug, as well as increased its stability and half-life in circulation (Blanco et al., 2011; Pearce et al., 2012). In addition, nanoparticles encapsulated with chemotherapy agents are able to avoid multi-drug resistant efflux pumps expressed on the surface of most tumor cells (Blanco et al., 2011; Pearce et al., 2012). The first nanoparticle-drug approved by the FDA (1995) was “Doxil.” Doxil is a liposome with an approximate size of 100 nm in diameter which encapsulates the chemotherapeutic drug doxorubicin (Barenholz). To improve tumor bioavailability the liposome was modified to contain a small amount of polyethylene glycol (PEG)-lipid to reduce the clearance of the nanoparticles from the blood, and increase the plasma half-life of doxorubicin. Indeed, encapsulation of doxorubicin within the liposome was shown to significantly alter its pharmacokinetic and pharmacodynamic properties leading to increased tumor uptake and anti-cancer activity, along with a reduction in systemic off-target toxicity (Barenholz, 2012). Today Doxil is used to treat a number of different solid tumors, including platinum-resistant ovarian cancer (Leamon et al., 2013).

Over the recent years, there have been attempts to develop nanoparticle formulations of the chemotherapeutic agent gemcitabine. Indeed, gemcitabine has long been the first-line treatment for patients with unresectable locally advanced or metastatic pancreatic cancer (Vincent et al., 2011). However, despite its use as a first-line treatment, patient survival has only been extended by 6–12 weeks (Hidalgo, 2010). Hence, in an attempt to improve the delivery of gemcitabine to pancreatic tumors as well as overcome some of the acquired gemcitabine-resistant mechanisms in pancreatic cancer cells nanoparticles have been used as a delivery vehicle. Recently, Wonganan et al. (2013), Zhu et al. (2013) demonstrated the potential of a nanoparticle encapsulated with the pro-drug of gemcitabine [4-(N) stearylgemcitabine] (GemC18) to overcome resistance associated with ribonucleotide reductase subunit M1 overexpression in pancreatic cancer cells. Interestingly, the authors showed that encapsulation of the gemcitabine pro-drug into the nanoparticle allowed for a different mode of entry into the cell which allowed the gemcitabine to be hydrolyzed more efficiently to its active form compared to the pro-drug alone (Wonganan et al., 2013; Zhu et al., 2013). In another example, Lee et al. (2013) engineered IONPs to express a uPAR-targeted moiety on their surface. Both pancreatic tumor cells and the blood vessels within the tumor stroma have high amounts of the uPAR receptor (Harvey et al., 2003). The magnetic IONPs also had gemcitabine attached via a lysosomally cleavable tetrapeptide linker (Lee et al., 2013). Importantly, systemic delivery of the nanoparticle-gemcitabine complex significantly inhibited the growth of orthotopically xenografted pancreatic tumors in mice and allowed for the detection of residual tumors following treatment using MRI (Lee et al., 2013). More recently, there has also been interest in developing nanotechnology to improve the delivery and efficacy of other chemotherapeutic agents which can be delivered in combination with gemcitabine or used as second-line treatment for pancreatic cancer. For example, Cabral et al. (2013) synthesized micelle nanoparticles which were able to self-assemble with the chemotherapeutic drug oxaliplatin. These nanoparticles were designed to gradually release their contents over time when only exposed to the tumor microenvironment. Importantly, the authors demonstrated that repeated systemic administration of the drug-loaded nanoparticles was able to significantly reduce tumor growth as well as the incidence of metastases in a clinically relevant transgenic mouse model of pancreatic cancer. Therefore, this therapy may be beneficial for treatment of patients with early-stage pancreatic cancer so as to prevent or retard the development of metastases.

Another nanoparticle-bound chemotherapy agent which has generated significant interest is albumin-bound paclitaxel known as Nab-paclitaxel or Abraxane® (Abraxis Bioscience). A recent phase I/II trial for pancreatic cancer demonstrated the maximum-tolerated dose for Nab-paclitaxel in combination with gemcitabine (Von Hoff et al., 2011). The authors also reported an improved overall survival in patients treated with nab-paclitaxel plus gemcitabine (12.2 median months of overall survival) compared to gemcitabine alone. Moreover, Nab-paclitaxel alone and in combination with gemcitabine was shown to deplete pancreatic stroma in pancreatic cancer xenograft mouse models (Von Hoff et al., 2011). Importantly, the depletion of stroma led to a 2.8 fold increase in the intratumoral concentration of gemcitabine (Von Hoff et al., 2011). A recent phase III MPACT (Metastatic Pancreatic Adenocarcinoma Clinical Trial) trial also showed that the addition of Nab-paclitaxel with gemcitabine was not only able to significantly improve the median survival of metastatic pancreatic cancer patients (8.5 months) when compared to gemcitabine treated only arm (6.7 months), but also significantly reduced toxicities (neuropathy and neutropenia) commonly associated with the cremaphor formulation used to dissolve paclitaxel thereby, allowing for a higher paclitaxel dose to be delivered (Ma and Hidalgo, 2013; Von Hoff et al., 2013). The authors also reported an increase in peripheral neuropathy and myelosuppression. However, these side-effects appeared to be reversible (Von Hoff et al., 2013).

Finally, nanotechnology had also been used to improve drug kinetics and tumor bioavailability, of a therapeutic agent by directly targeting and ablating the tumor stroma. Indeed, Provenzano et al. (2012) showed that the desmoplastic reaction surrounding pancreatic tumor cells generated very high amounts of interstitial fluid pressure along with the induction of vascular collapse (Provenzano et al., 2012). Systemic administration of PEGylated human recombinant PH20 hyaluronidase (PEGPH20) [an enzymatic agent that targets a critical component of the desmoplastic stroma in pancreatic cancer known as hyaluronic acid (HA)] (Provenzano et al., 2012) in a murine pancreatic cancer model, produced a marked decrease in tumor stroma which correlated to a rapid and significant decrease in interstitial tumor pressure and increased tumor blood vessel lumen diameter. Furthermore, when delivered in combination with gemcitabine there was a strong anti-tumor effect compared to gemcitabine alone (Provenzano et al., 2012). Collectively, these studies highlight the advances in nanotechnology as highly effective carriers for the passive and active delivery of chemotherapy drugs to tumor cells in pre-clinical and clinical settings. In the future, nanotechnology which can therapeutically target the tumor stroma to enable stroma depletion and tumor vascular normalization as
well as deliver therapeutics directly to the tumor cells may become a highly effective novel treatment strategy for pancreatic cancer.

**NANOPARTICLES AS DELIVERY VEHICLES FOR RNA INTERFERENCE INHIBITORS**

RNA interference (RNAi) based therapeutics are emerging as an innovative and promising alternative over conventional systemic treatments in terms of specificity, toxicity, and overcoming multiple drug resistance. RNAi is an endogenous gene-silencing mechanism that can cause the degradation of any mRNA, once the RNA target sequence is known (Rana, 2007). In particular, short-interfering RNAs (siRNA) have gained attention due to their ability to potently silence target gene expression both in vitro and in vivo (Rana, 2007). siRNAs are processed double stranded RNAs approximately 21 nucleotides in length, and are involved in post-transcriptional gene silencing toward targeted mRNAs. The RNA-induced silencing complex (RISC) located within the cytoplasm of the cell act as a guide for the cleavage of mRNAs bearing a complementary sequence to the siRNA (Rana, 2007). Once activated RISC can be recycled multiple times to cleave additional mRNA targets (Rana, 2007). Importantly, the introduction of chemically synthesized siRNAs into the cell can activate this naturally-occurring mechanism, which can be harnessed as a powerful gene therapy to suppress specific genes associated with human disease, including cancer. However, delivery of siRNA to a target cell remains a challenge. The major limitation of in vivo siRNA delivery is its instability and vulnerability to degradation in serum and inability to readily enter a cell due to its high anionic (negative) charge (Baigude and Rana, 2009). Nanoparticles in the form of liposomes, lipid polymers, and dendrimers have been developed over the last decade to act as highly effective siRNA delivery vehicles (Figure 1) (Zimmermann et al., 2006; Baigude et al., 2007; Davis et al., 2010; Su et al., 2011). Indeed, nanoparticles can be designed to self-assemble with siRNA and protect it from serum degradation and elimination from the body (Schroeder et al., 2010). Moreover, nanoparticles can be tailored to possess multifunctional components to allow for targeted siRNA delivery and efficient entry to a specific cell type (Schroeder et al., 2010). The therapeutic potential of nanoparticle-siRNA complexes to treat human disease was first reported by Zimmermann et al. (2006). In this study, the authors used modified nanoparticles (liposomes) known as Stable Nucleic Acid Lipid Particles (SNALP) to deliver siRNA targeting apolipoprotein B (apo B) (a protein involved in regulating cholesterol metabolism) to non-human primates. A single systemic administration of low clinically relevant amounts of

![Nanoparticle-siRNA delivery vehicles](image)
SNALP-siRNA complexes caused a significant reduction in apo B mRNA and protein expression as well as a reduction in total cholesterol (Zimmermann et al., 2006). The nanoparticle-siRNA complexes were non-toxic and provided sustained knockdown of apo B for up to 11 days (Zimmermann et al., 2006). SNALPs have also been used to deliver siRNA to potently silence a gene involved in promoting aggressive tumor growth in an orthotopic mouse model of liver cancer (Judge et al., 2009). Collectively, these studies highlight the potential of using nanoparticle-siRNA complexes as novel therapeutics to treat human disease in the clinic.

Recently there has been intense research effort into the development of nanoparticles which contain targeting moieties (antibodies, peptides) conjugated to their surface to deliver siRNA specifically to tumor cells (Schoeders et al., 2010). Common examples of cell surface targets used for delivery to tumor cells include: transferrin receptor (TfR), folate receptor, and Arginine-Glycine-Aspartic ligands. It is to be noted that these cell surface receptors have been reported to be highly expressed in a number of different tumor types (Allen, 2006; Daniels et al., 2006; Basile et al., 2012). For instance, Piroolio et al. (2007) used nanoparticles (liposomes) which contained a TfR-antibody on their surface to complex siRNA against HER-2. Systemic delivery of the TfR-targeting nanoparticles resulted in more siRNA being delivered to pancreatic tumors in a murine xenograft model compared to nanoparticles without the TfR antibody (Piroolio et al., 2007). Moreover, delivery of the nanoparticle-siRNA complexes were able to silence HER-2 expression in the pancreatic tumor cells and increase sensitivity to gemcitabine (Piroolio et al., 2007). Studies have also taken interest in targeting nanoparticles carrying siRNA to the stroma of the pancreas. Of note, Ishiwatari et al. (2013) were able to target activated PSCs using vitamin A-coupled nanoparticles (liposomes) loaded with siRNA against collagen-specific chaperone protein gp46 (VA-lip-siRNAgp46) (Ishiwatari et al., 2013). VA-lip-siRNAgp46 complexes were actively taken up by PSCs and subsequent silencing of gp46 expression correlated to a significant decrease in PSC collagen secretion, which in turn resolved pancreatic fibrosis in a rat model of chronic pancreatitis (Ishiwatari et al., 2013). These promising findings point toward the therapeutic potential of nanoparticle-siRNA complexes in ablating the desmoplasia in pancreatic cancer.

Finally, siRNA-nanoparticle therapy based human clinical trials are underway to evaluate the safety and efficacy of systemic siRNA delivery using modified nanoparticles for the treatment of cancer (Burnett et al., 2011; Davidson and McJay, 2011). Most notably, a phase I clinical trial involving systemic siRNA administration to patients with solid tumors using a TfR-guided cyclodextrin-nanoparticle (CALAA-01) showed great promise (Davis et al., 2010). CALAA-01 is a nanoparticle-siRNA complex that was used to silence anti-ribonucleotide reductase (RRM2) in patients with metastatic melanoma. Results from this study showed that systemic administration of CALAA-01 reduced RRM2 mRNA and protein levels in tumor tissue collected via biopsy after administration (Davis et al., 2010). Another landmark phase I study recently reported the use of nanoparticles to simultaneously deliver two individual siRNAs targeting vascular endothelial growth factor (VEGF) and kinesin spindle protein (KSP) (Tabernero et al., 2013). The nanoparticle-siRNA complex later renamed ALN-VSP was administered bi-weekly (intravenously) and was found to be safe and well tolerated even up to 23 months (Tabernero et al., 2013). Importantly, results also showed disease control lasting more than 6 months and a complete regression of liver metastases in a patient with endometrial cancer (Tabernero et al., 2013).

CONCLUDING REMARKS

New strategies are needed to reduce the overall dismal prognosis and increase survival of pancreatic cancer patients. Nanomedicines offer great potential with benefits for diagnostics, imaging, and therapeutics. As our knowledge of pancreatic cancer becomes more complete, it is increasingly important for clinicians, biologists, and biochemical engineers to integrate novel ideas for the treatment of pancreatic cancer. Developing nanoparticle therapies based on the unique tumor microenvironment is crucial to the delivery of clinically relevant amounts of active therapeutics to the tumor site while by-passing various biological barriers. As highlighted in this review nanoparticles could be used to target tumor elements and stromal elements of pancreatic cancer. In summary, nanotechnology will have an important role in realizing the goal for early detection and personalized pancreatic cancer treatment for patients at different stages of disease, avoiding unnecessary toxicities associated with current treatment regimes and increasing patient survival.

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