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Chapter 5

Gd-Doped Superparamagnetic Magnetite Nanoparticles for Potential Cancer Theranostics

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Additional information is available at the end of the chapter

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Abstract

Nanotechnology has facilitated the applications of a class of nanomaterials called superparamagnetic iron oxide nanoparticles (SPIONs) in cancer theranostics. This is a new discipline in biomedicine that combines therapy and diagnosis in one platform. The multifunctional SPIONs, which are capable of detecting, visualizing, and destroying the neoplastic cells with fewer side effects than the conventional therapies, are reviewed in this chapter for theranostic applications. The chapter summarizes the design parameters such as size, shape, coating, and target ligand functionalization of SPIONs, which enhance their ability to diagnose and treat cancer. The review discusses the methods of synthesizing SPIONs, their structural, morphological, and magnetic properties that are important for theranostics. The applications of SPIONs for drug delivery, magnetic resonance imaging, and magnetic hyperthermia therapy (MHT) are included. The results of our recent MHT study on Gd-doped SPION as a possible theranostic agent are highlighted. We have also discussed the challenges and outlook on the future research for theranostics in clinical settings.

Keywords: theranostics, Fe₃O₄ nanoparticles, MRI contrast agent, drug delivery, magnetic hyperthermia

1. Introduction

Nanomaterials, with the size of at least one dimension ranging from a few nanometers to about a hundred nanometers, having unique properties compared to their respective bulk materials, are of intense research interest because of their applications in various fields of science and technology. One of the major applications, among many of their potential applications, is in biomedicine as platform for effective diagnosis and therapy [1–3]. The multifunctionality of
these nanoparticles has recently led the biomedicine research in a new direction called “Theranostics” which is the integration of diagnostic imaging and therapeutic function into a single platform [4, 5]. Theranostic agents allow the combination of diagnosis, treatment, and follow-up of a disease and hence are expected to contribute to personalized medicine. Among many nanomaterials, magnetic nanoparticles (MNPs) have the potential to deliver imaging and therapeutic agents to a specific region in the body with an external magnetic field manipulation. This requires large magnetization for the MNP so that they could respond to externally applied magnetic fields at physiological temperatures. Superparamagnetic iron oxide nanoparticles (SPIONs), such as Fe$_3$O$_4$ and $\gamma$-Fe$_2$O$_3$ nanoparticles, exhibit relatively higher saturation magnetization with no magnetic hysteresis (zero remanence and coercivity) and fulfill other major requirements such as low toxicity, biocompatibility, and surface functionalization capabilities for theranostic applications. A number of SPIONs have undergone clinical trials and several formulations have been approved for clinical imaging and therapeutic applications [6]. A few examples are Lumiren for bowel imaging, Ferridex IV for liver and spleen imaging, Combidx for lymph node metastases imaging, and Ferumoxytol for iron deficiency therapy.

Furthermore, SPIONs can be multipurposely used for diagnosis such as magnetic resonance imaging (MRI) and for therapeutic functions such as targeted delivery of therapeutic agents, anticancer drugs, siRNA, and for magnetic hyperthermia (MHT) for cancer treatments. This makes SPION an ideal vehicle in the development of theranostic nanomedicine [7–9]. An example of strategy for using magnetic nanoparticles as a potential theranostic agent is illustrated in Figure 1.

In this chapter, we discuss the detailed background on magnetic properties of SPIONs and their synthesis methods and surface modification for cancer diagnosis and therapy. In addition, various applications of SPION ranging from MRI contrast agent to therapeutic-targeted drug delivery and MHT are discussed. We have also highlighted the results of our recent study on Gd-doped SPION as a possible theranostic agent. The remainder of the chapter focuses on the challenges and outlook on the future research for theranostics in clinical settings.

Figure 1. Schematic illustration of the therapeutic strategy using MNP. Functionalized MNPs accumulate in the tumor tissues via the drug delivery system. MNP can be used as a tool for cancer diagnosis by MRI or for magneto-impedance sensor. Hyperthermia can then be induced by alternating magnetic field exposure.
2. Magnetic properties of SPION

2.1. Background

MNP s have been studied for over 50 years now due to their potential application in many areas including biomedical sciences. As for the types of MNP, the major focus has been on iron oxide (Fe₃O₄), gold-coated iron oxide (Au-Fe₃O₄), metallic iron (Fe), and Fe-Co and Fe-Pt nanoparticles. In most cases, the particle size ranges from 1 to 100 nm exhibiting high surface-to-volume ratio. As a result, they offer higher surface area for interaction with foreign objects compared to larger particles. Many review articles have been written that focus on sensing, drug delivery, and hyperthermia properties of these nanoparticles [10–13]. The physical, chemical, and magnetic properties of MNP largely depend on synthesis method and their surface modification, and much progress has been made in this direction to MNP of varying sizes, shapes, composition, and core-shell designs [14–23].

The important magnetic parameters relevant to theranostic applications are saturation magnetization (Mₛ), remanent magnetization (Mᵣ), coercivity (Hcdc), Curie temperature (Tc), magnetic anisotropy energy density (K), and blocking temperature (Tb). These parameters are influenced by the material, size, shape, composition, and core-shell (functionalization) of the nanoparticles. Mₛ is the maximum value of magnetization of the material that can be achieved under the influence of an external magnetic field, Mᵣ is the remanent magnetization in the material after removing the external magnetic field, Hcdc is the strength of the reverse magnetic field needed to bring the remanent magnetization to zero, and K is the material property signifying the tendency of the magnetization to orient along a certain axis of the particle. As the volume (V) of the particles decreases, the magnetic anisotropy energy (KV) of the nanoparticle also decreases. If the particle size is reduced below a certain critical size, it becomes a single magnetic domain creating a giant spin called “superspin” leading to a large magnetic moment (~10,000 Bohr magneton) on each particle. The behavior of a collection of such noninteracting particles under an external magnetic field is determined by a competition between the magnetic anisotropy energy barrier (ΔE) and the thermal energy (k_BT) for magnetic moment reversal. Above a characteristic temperature called the blocking temperature, T_b, their behavior is very similar to that of a paramagnetic material and described as “superparamagnetism.” The underlying physics of superparamagnetism is founded on the activation law for the relaxation time τ of the net magnetization of the particle given by τ = τ₀ exp (ΔE/k_BT), where τ₀ is of the order of 10⁻¹⁰–10⁻¹² s [24].

2.2. Effect of size, shape, and composition

Magnetic properties of materials, such as susceptibility, coercivity, and saturation magnetization, depend on the structure, size, shape, and composition, and can be altered to manipulate the magnetic properties. Particle size plays an important role in many magnetic biomedical applications such as magnetic hyperthermia and drug delivery, where the size used lies in the nanometer regime. The MNPs often contain a layer of disordered spins on the surface of the particle leading to reduction in their Mₛ compared to the corresponding bulk material. A relation between the Mₛ and the size of the nanoparticle is given by [25]
where $r$ is the radius of the nanoparticle, $d$ is the layer thickness of the disordered spins, and $M_{sb}$ is the saturation of the bulk material. Recent studies have shown that the functionalization of MNP can reduce the thickness of the surface-disordered spin layer [26].

Although the effect of shape of MNP on their magnetic properties is not extensively studied, a few investigations have been reported in the literature on ferrite nanocubes, maghemite nanorods, NiFe wires, cobalt nanodiscs, tetrapods, and Au-MnO nanoflowers showing a strong dependence of $M_s$ on the shapes of the nanoparticles [27–36]. Higher $M_s$ values have been observed for the cubic MNP compared to the spherical MNP of the same size [37]. Also, cubic Fe$_3$O$_4$ nanoparticles have been found to exhibit higher $T_c$ compared to spherical Fe$_3$O$_4$ nanoparticles [38], and the amount of disordered spins to be less (4%) in the former and more (8%) in the latter [39].

Magnetic properties of widely used magnetite (Fe$_3$O$_4$) with its spinel structure of $[\text{Fe}^{3+}]_{A}[\text{Fe}^{3+}\text{Fe}^{2+}]_{B}\text{O}_4$ can be changed if other magnetic atoms such as Ni, Co, Mn, and so on are substituted at the tetrahedral A or octahedral B sites of the spinel structure. This flexibility of creating mixed ferrites is useful in tuning the magnetic properties for hyperthermia applications. There have been numerous studies investigating the interdependence of magnetic properties and the composition. The method of preparation, concentration and nature of dopants, and postsynthesis processes have shown to profoundly affect the magnetic properties. A study [40] compared the magnetization among the four spinels of FeFe$_2$O$_4$, MnFe$_2$O$_4$, CoFe$_2$O$_4$, and NiFe$_2$O$_4$ for the same size of 12 nm and found the highest magnetization for MnFe$_2$O$_4$. In another study with Y$_3$Fe$_{5-x}$Al$_x$O$_{12}$ for $x$ varying between 0 and 2, the Curie temperature changed from 40 to 280°C [41]. With increasing Al, Fe$^{3+}$ cations occupied the tetrahedral sites and some of the octahedral sites of Fe$^{2+}$ were replaced by nonmagnetic Al$^{3+}$ cations, which reduced the magnitude of $M_s$. The $T_c$ value reached the room temperature for $x$ value between 1.5 and 1.8. The variation in composition affects not only the magnitude of $M_s$ but also the coercivity. The tailoring of ferromagnetic to paramagnetic phase transition temperature is particularly very useful in hyperthermia application to turn off undesirable heating beyond the required temperature.

3. SPION synthesis and surface modification

Over the past decades, many efficient synthesis methods have been developed to produce the size/shape controlled, stable, biocompatible, and monodispersed iron oxide nanoparticles [42–45]. The most common methods include coprecipitation [46, 47], thermal decomposition [48], hydrothermal synthesis [49, 50], microemulsion [51], and sonochemical [52] synthesis. Thermal decomposition technique involves decomposition of organo-metallic iron precursors in organic solvents at higher temperatures. Although the method can produce high-quality monodisperse particles because of separate nucleation and growth processes, it is a complicated synthesis method and
produces hydrophobic nanoparticles that cannot be directly used for bio-applications without laborious postsynthesis processes, which may result in aggregation and loss of magnetic properties. The most commonly used technique is the coprecipitation method, which is a cost-effective and a facile synthesis method. However, this method produces Fe₃O₄ nanoparticles with wide particle size distribution due to lack of control over hydrolysis reactions of the iron precursors, and the nucleation and growth steps leading to particles with a range of superparamagnetic-blocking temperature. The other common, recently developed, method is the hydrothermal synthesis, which generates nanoparticles with excellent crystallinity with controllable size and shape in aqueous phase. The properties of the nanoparticles can vary with the synthesis method due to the differences in cationic distribution and vacancies, spin canting, or surface contribution.

In the design of magnetic nanoparticles for theranostic applications, surface modification plays an important role in providing colloidal stability and biocompatibility. The stable colloidal suspensions of surfactant-coated SPION are called “ferrofluids” which are magnetizable and remain as liquids in the presence of magnetic fields and in biological media. Stabilization of the ferrofluid occurs in the presence of one or both of the two repulsive forces (see Figure 2). The electrostatic repulsion can be understood through the knowledge of the diffusion potential and mainly depends on the ionic strength and the pH of the solution. The steric force is difficult to predict or quantify and mostly depends on the weight and the density of the polymer used for the coating.

In order to achieve biocompatibility, the coating should prevent any toxic ion leakage from magnetic core into the biological environment as well as shielding the magnetic core from oxidation and corrosion. When nanoparticles are injected into the body during in vivo applications, they are often recognized by reticuloendothelial system (RES) that eliminates any foreign substance from blood stream [53]. In this process, nanoparticles are rapidly attacked by the plasma proteins from RES and shuttled out of circulation to the liver, spleen, or kidney.
which are then cleared out from the body. Also, this RES accumulation often causes toxicity issues as well. The specific surface coatings can prevent the adsorption of these proteins, increasing the circulation time in blood, hence maximizing the possibility to reach target tissues [54]. For instance, it is well known that coating of hydrophilic polymers, mainly polyethylene glycol (PEG), on the nanoparticles reduces nonspecific binding of the proteins resulting in stealth behavior. In addition to the stabilization and enhanced biocompatibility, these protecting shells also provide a platform for further functionalization such as the addition of specific targeting ligands, dyes, or therapeutic agents.

Over the years, researchers have developed various surface modification strategies composed of grafting of or coating with both organic and inorganic materials. Organic molecules include small organic molecules, macromolecules or polymer and biological molecules. They provide various highly reactive functional groups such as carboxyl groups, aldehyde groups, and amino groups. Polymer-coating materials can be classified into synthetic and natural, and some commonly used polymers are listed in Table 1 along with their advantages.

The surface coating could affect the magnetic properties of SPION. Many studies have reported the effect of the surfactants on the magnetic properties [55–60]. Yuan et al. [58] investigated the effect of surfactant on magnetic properties using commercially available aqueous nanoparticle suspensions, FluidMAG-Amine, FluidMAG-UC/A, and FluidMAG-CMX, in parallel with oleic acid-covered particles suspended in hexane and heptane. Their results reveal the reduction of magnetic phase in nanoparticles, which varies with different coatings as well as with solvents. The reduction in magnetization with different coatings was attributed to the different degree of surface spin disorder.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Advantages and applications</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natural</td>
<td>Dextran Stability, biocompatibility, enables optimum polar interactions with iron oxide surfaces, and enhances the blood circulation time</td>
<td>[61–66]</td>
</tr>
<tr>
<td>Starch</td>
<td>Improves the biocompatibility, good for MRI, and drug target delivery</td>
<td>[67, 68]</td>
</tr>
<tr>
<td>Chitosan</td>
<td>Biocompatible and hydrophilic large abundance in nature, biocompatibility, and ease of functionalization, widely used as nonviral gene delivery system</td>
<td>[69–72]</td>
</tr>
<tr>
<td>Synthetic</td>
<td>Poly(ethylene-glycol) (PEG) Enhances the hydrophilicity and water-solubility, improves the biocompatibility, blood circulation times, and internalization efficiency of the nanoparticles. Used in target-specific cell labeling, magnetic hyperthermia, targeted drug delivery</td>
<td>[73–75]</td>
</tr>
<tr>
<td>Alginate</td>
<td>Improves the stability and biocompatibility. Used in drug delivery applications</td>
<td>[76–78]</td>
</tr>
<tr>
<td>Poly-N-isopropylacrylamide (PNIPAM)</td>
<td>Generally used as thermosensitive drug delivery and cell separation</td>
<td>[79–82]</td>
</tr>
<tr>
<td>Polyethylene-imine (PEI)</td>
<td>Ability to complex with DNA, guide intracellular trafficking of their cargo into the nucleus, used for gene delivery cell transfection with either DNA or siRNA nucleotides</td>
<td>[83–86]</td>
</tr>
</tbody>
</table>

Table 1. Commonly studied organic polymers and their advantages.
4. SPION for drug delivery

Over the last two decades, MNPs have been increasingly exploited as platforms for the transport of therapeutics including drugs and genes [46, 87, 88]. In magnetic drug delivery, a drug or a therapeutic reagent is conjugated to the nanoparticle and introduced in the body, and concentrated in the target area by means of a magnetic field gradient (using an internally implanted permanent magnet or an externally applied field) [89]. Even though magnetic drug delivery shows a great promise in cancer treatment avoiding the side effects of conventional chemotherapy, the designing and fabrication of an efficient nanoparticle-based drug delivery system is still a challenge. Using a targeting ligand, the targeting specificity can be enhanced. These anticancer drugs carried by the nanoparticles can then be released at the tumor site via enzymatic activity, or via changes in the physiological conditions such as temperature and pH. Drug release can also be magnetically triggered from the drug-conjugated magnetic nanoparticles [89–91]. For example, Hayashi et al. [92] reports a study done on superparamagnetic iron oxide nanoparticles conjugated with folic acid (well known as a targeting ligand for breast cancer cells), β-Cyclodextrin (which acts as drug container), and Tamoxifen (anticancer drug). Using an AC magnetic field, heat is generated which triggers drug release—a behavior that is controlled by switching the high-frequency magnetic field on and off. This is capable of performing drug delivery and hyperthermia simultaneously. Among various other anticancer drugs, Doxorubicin (Dox) is widely used as a model drug. There are several methods that can be used to load Dox into nanoparticles, such as by adsorption onto nanocarrier inorganic core [93–95], by diffusion [95, 96], or entrapment [97, 98] in the coating materials and by chemical bonds [99, 100] with the coating of the nanocarrier. Several modifications including surface functionalization of these SPIONs with Dox have been conducted over the last few years to investigate their efficacy [101]. Previous studies have reported that PEG-functionalized porous silica shell onto Dox-conjugated Fe₃O₄ nanoparticle cores [102], PAMAM (Poly(amidoamine))-coated Fe₃O₄ nanoparticles-Dox complex [103], and Dox-loaded Fe₃O₄ nanoparticles modified with PLGA-PEG copolymers [104] could potentially be very promising in therapeutic cancer treatment. However, most of the Dox-SPION–based drug delivery studies have been focused on human breast cancer cells.

In our recently published work [105], we have developed a novel drug delivery platform based on Fe₃O₄ nanoparticles as a vehicle for an anticancer drug (Dox), attached to a model dye (FITC) for their precise tracking and investigated their incorporation into the human pancreatic cancer cell line (MIA PaCa-2) for specific drug targeting. Existing EDC/NHS technique was employed for this dual drug/dye conjugation. This unique drug-dye dual conjugation of SPION after penetration through the cell membrane shows a steady release of Dox into the nucleus of the malignant cells. Our studies demonstrate that the association of Dox onto the surface of nanoparticles enhances its penetration into the cancer cells as compared to the unconjugated drug as shown in the subsequent text (Figure 3). In addition to the rapid uptake of these SPIONs by live cells, our results also suggest that upon entering the cells, Dox is cleaved from the conjugation, which might be due to the enzymatic reactions that occur within the cells, and tends to accumulate in the nuclei fulfilling the major requirement for an effective therapeutic system.
5. SPION for cancer diagnosis using MRI

Magnetic resonance imaging with its high spatial resolution has been a preferred method of imaging and diagnosing a disease. It is a noninvasive medical diagnostic tool that monitors the change in magnetization of hydrogen-protons in water molecules contained in a tissue when placed in a magnetic field and exposed to a pulse of radio frequency electromagnetic waves. The mapping of the magnetization provides an image of the organ due to the fact that protons in different tissues, with varying water concentration, respond differently. Contrast agents have been used to enhance the images as they affect the behavior of the protons in their vicinity leading to sharper images. Contrast agents used in MRI are divided into two categories: $T_1$ and $T_2$.
$T_2$ contrast agents based on their effect on the magnetic relaxation processes of the protons [106]. Most commonly used $T_1$ contrast agents are paramagnetic compounds that are composed of metal ions of Gd$^{3+}$ or Mn$^{2+}$ and a chelating ligand, such as diethylene triamine penta-acetic acid, DTPA [106, 107]. The chelate prevents the metal ion from binding to chelates in the body making the paramagnetic ion less toxic. $T_1$ contrast agents mainly reduce the longitudinal relaxation time ($T_1$) which is due to energy exchange between the spins and surrounding lattice (spin-lattice relaxation) and result in a brighter signal. $T_2$ contrast agents, consisting of superparamagnetic nanoparticles such as Fe$_3$O$_4$, have a strong effect on the transverse relaxation time ($T_2$). In an external magnetic field, nanoparticles are magnetized and generate induced magnetic fields locally. These induced fields perturb the magnetic relaxation processes of the protons in the water molecules decreasing the $T_2$ relaxation time, which results in darkening of MR images.

There are various research studies conducted on enhancing the MRI signal for cancer detection using SPIONs as $T_2$ contrast agents [108–110]. The efficiency of SPIONs as $T_2$ contrast agents mainly depends on their physicochemical properties, particularly their size and surface chemistry. Stephen et al. [111] report the correlation between particle size and $T_2$ relaxation. Their study shows that a decrease in particle size leads to reduction in saturation magnetization, which in turn reduces the $T_2$ relaxation capabilities of SPIONs. There are studies which show the effect of shape on relaxivity. For example, Zhen et al. [112] reported that cubic Fe$_3$O$_4$ MNP showed four times smaller relaxation time and thus better image contrast compared to the spherical Fe$_3$O$_4$. In another study, octapod Fe$_3$O$_4$ nanoparticles with an edge length of 30 nm show a smaller value of $T_2$ compared to 16-nm spherical Fe$_3$O$_4$ nanoparticles possessing a similar $M_s$ [37]. The studies by Park et al. [113] report a decrease in relaxivity as PEG molecular weight increases, indicating that the thickness of PEG coating at the particle surface affects $T_2$ relaxivity.

When using SPIONs as contrast agents for MRI, it is crucial that they are captured into the cells efficiently upon exposure. Some approaches include introducing peptides [114], antibodies [115], and polymers [116] onto or surrounding magnetic nanoparticles to improve the target specificity. For example, Jun et al. [117] have successfully synthesized superparamagnetic iron oxide nanoparticles of 9-nm size as magnetic probes for the in vivo detection of cancer cells implanted in a mouse. In their research work, 2,3-dimercaptosuccinic acid (DMSA) ligand is attached to the nanoparticles surface to obtain hydrophilic nanoparticles and the nanoparticles are further conjugated with the cancer-targeting antibody, Herceptin. The specific binding properties of Herceptin against a HER2/neu receptor overexpressed from breast cancer cells lead to the successful detection of breast cancer cells (SK-BR-3).

Even though both $T_1$ and $T_2$ mapping are powerful techniques, single-mode contrast agents are not always sufficient in modern diagnosis as they have certain drawbacks and limitations [118]. For example, the dark contrast produced by $T_2$ agents can also be generated from adjacent bones or vascular or there can be susceptibility artifacts due to the sharp change in magnetic field at the surrounding contrast agent. Also, Gd-chelates ($T_1$ agent) have high mobility which shorten their presence in the vascular system and raise possible toxicity issues. Thus, there is a growing interest in developing complementary $T_1$-$T_2$ dual-modal contrast agents, combining the advantages of...
positive and negative contrasts to obtain high sensitivity and biocompatibility for improved diagnosis [119]. Two different approaches of integrating $T_1$ and $T_2$ species have been reported recently [118]. One is constructed by labeling $T_1$-signaling elements (Gd species) on magnetic nanoparticles. In the study done by Bae et al. [120], Gd-DTPA, a representative Gd chelate-based $T_1$ MRI contrast agent, is covalently attached to dopamine-coated iron oxide nanoparticles. Their results demonstrated that the composite not only had the ability to improve surrounding water proton signals on the $T_1$-weighted image but also could induce significant signal reduction on the $T_2$-weighted image. In another study reported by Santra et al. [121], Gd-DTPA is encapsulated within the poly (acrylic acid) (PAA) polymer-coated SPION (IO-PAA) conjugated to folic acid, which acts as the targeting ligand for breast cancer cells (HeLa cells). When nanoprobes are internalized within the cells, which is acidic, composite magnetic nanoprobe degrades resulting in an intracellular release of Gd-DTPA complex with subsequent $T_1$ activation, which can be seen by MRI. Authors claim that this $T_1$ nano-agent could be used for the detection of acidic tumors. The other type of conjugated system consists of $T_1$ paramagnetic elements embedded into $T_2$ magnetic nanoparticles. For example, Zhou et al. [122] have synthesized Gd$_2$O$_3$-embedded iron oxide nanoparticles with an overall size of 14 nm which can act as a $T_1$-$T_2$ mutually enhanced dual-modal contrast agent for MR imaging of liver and hepatic tumor detection with great accuracy in mice. Xiao et al. [123] have prepared PEGylated, Gd-doped iron oxide nanoparticles which is applicable as a $T_1$-$T_2$ dual-modal MRI contrast agent. Their in vivo MRI results demonstrated the simultaneous contrast enhancements in $T_1$- and $T_2$-weighted MR images toward the glioma-bearing mice.

6. SPION for cancer therapy using magnetic hyperthermia

Magnetic hyperthermia is the transformation of electromagnetic energy from an external alternating magnetic field into heat using MNP. Magnetic nanoparticles serve as the nano-heat centers producing heat by relaxation losses, thereby heating the tissue. The main goal of an effective cancer treatment is to kill the malignant cells with the least of damage to normal cells. As MHT can be used for heating small regions selectively, it offers the potential for being highly selective and noninvasive technique for therapeutic treatment of cancers, and consequently it has advantage over other treatment such as chemotherapy and radiation therapy. It is known that reduced blood flow in tumor causes the lack of oxygen in tumor site which leads to the formation of lactic acid making the cells more acidic [124]. The acidic cells are more sensitive to temperature, have lower thermal resistance than normal cells, and the decreased blood flow in the tumor limit their ability to dissipate heat. As a result, cancer cells can be damaged or killed by increasing the local temperature to the range of 42–46°C with little detriment to healthy cells.

The idea of utilizing SPION for hyperthermia was first proposed by Gilchrist et al. [125] in 1950s and since then many types of MNP are being investigated for this purpose. MNPs have the advantage of being guided and localized specifically at a tumor site by external magnetic fields and can also be directed to the cancer cells by tagging a ligand, such as an antibody or a peptide, without reducing its efficiency. For example, Fabio et al. [126] have reported that the conjugation of folate receptors enhances the targeting for magnetic hyperthermia in solid
tumors. Magnetite (Fe$_3$O$_4$) and maghemite ($\gamma$-Fe$_2$O$_3$) have been extensively studied and are promising candidates due to their biocompatibility and relative ease for functionalization. Additionally, iron oxide nanoparticles doped with other magnetic dopants such as Co, Mn, and Ni [127–129] are under investigation to achieve a high heating efficiency by tuning the magnetic anisotropy and saturation magnetization of the material. In addition, many recent findings show that multicore nanoparticles possess a higher heating power than the single-core particles [130], and may offer an advantage. However, among numerous complications, with a high Curie temperature of Fe$_3$O$_4$, 850 K, and $\gamma$-Fe$_2$O$_3$, 750 K, overheating is one of the drawbacks of utilizing these nanoparticles, and as a solution, those complex magnetic oxides with low Curie temperature are being investigated [131–133].

Specific absorption rate (SAR) is a measure of efficiency of heat generation. The SAR value can be estimated by measuring the temperature change in the ferrofluid samples upon exposure to an AC magnetic field following the equation [134]:

$$\text{SAR}(T) = \frac{M_{\text{sample}}}{m_{\text{Fe}_3\text{O}_4}} \left( \frac{\Delta T}{\Delta t} \right)_T$$

Here, $M_{\text{sample}}$ is the mass of the sample, $m_{\text{Fe}_3\text{O}_4}$ is the mass of Fe$_3$O$_4$ nanoparticles in the sample, $C$ is the specific heat capacity of the sample, and $\left( \frac{\Delta T}{\Delta t} \right)_T$ is the time rate of change of temperature at $T$ obtained from the slope of the time-dependent temperature data. SAR depends on magnetic properties of the particles such as $M_o$ anisotropy constant $K$, particle size distribution ($\sigma$), magnetic dipolar interactions, and the rheological properties of the target medium. An ensemble of poly-disperse particles is usually described by a log-normal distribution function,

$$f(D) = \frac{1}{\sqrt{2\pi}\sigma D} \exp\left\{ -\frac{[\ln(D/D_0)]^2}{2\sigma^2} \right\}$$

Here, $D_0$ is the most probable particle diameter and $\sigma$ is the width of the distribution. The temperature-dependent average power dissipation in the sample is expressed as [135]

$$\overline{P}(T) = \int_0^{\infty} \frac{\mu_o \chi_o H_0^2 \omega}{2} \frac{\omega \tau_{\text{eff}}}{1 + (\omega \tau_{\text{eff}})^2} f(D) d(D)$$

where $H_0$ and $\omega$ are the amplitude and angular frequency of the applied AC magnetic field, $\mu_o$ is the vacuum permeability, $\tau_{\text{eff}}$ is the effective relaxation time involving Néel relaxation and the Brownian relaxation times, and $\chi_o$ is equilibrium susceptibility. $\tau_{\text{eff}}$ is defined as $\frac{1}{\tau''} = \frac{1}{\tau_N} + \frac{1}{\tau_B}$ where $\tau_B = \frac{4\eta g_k}{R_c V_m}$ and $\tau_N = \frac{\mu_o^2}{2} \frac{\chi'}{T_0} \exp\left( \frac{\chi'}{2T} \right)$, with $\chi'$ are the Néel and Brownian relaxation times, $\eta$ is the viscosity of the suspension, $R_c$ is the hydrodynamic radius of the coated nanoparticle, $V_m$ is the magnetic volume of the nanoparticles, and $\tau_o = 10^{-9}$ s. $\chi_o$ is given by $\chi_o = \chi_i \left( \coth \xi - \frac{1}{\xi} \right)$, where $\chi_i = \frac{2M_o H_0 V_m}{k_B T}$ is the initial susceptibility, and $\xi = \frac{2M_o H_0 V_m}{k_B T}$, with $M_d$ being the domain magnetization of the nanoparticle and $\phi$ the volume fraction of the magnetic nanoparticles in the ferrofluid. SAR in units of W/g is obtained using Eq. (4) as $\overline{P}(T)/m_{\text{Fe}_3\text{O}_4}$, where $m_{\text{Fe}_3\text{O}_4}$ is the mass of Fe$_3$O$_4$ nanoparticles in ferrofluids.
MHT investigations are often done on colloidal suspensions of surface-coated MNP, called ferrofluids. It is often necessary to coat the MNP using a biocompatible polymer to avoid direct contact with the tissue and to reduce the particle aggregation. Ferrofluid preparations frequently yield a mixture of isolated nanoparticles and nanoclusters [136] with varying degree of magnetic dipole-dipole interactions present in ferrofluids. It has been shown that the dipolar interactions among the MNP affect the SAR value drastically and can be exploited to optimize SAR [136–139]. A mean-field approximation method has been used to account for effects of interactions on SAR for a collection of monodisperse MNP [138, 139], and we have used this approach to explain the very different observed SAR values for similar size particles prepared by two different methods of preparation [140].

Since the surfactant influences the magnetic properties as well as the degree of interactions in the ferrofluid, a useful approach for improving the magnetic hyperthermia performance is to optimize the surface coating to maximize the SAR. Currently, there are conflicting SAR values obtained for a certain sized nanoparticle making it difficult to evaluate the exact contributions of surface coating on the SAR. According to Mohammad et al. [141], it is found that inorganic coatings improve the SAR value and the gold coating retains the superparamagnetic fraction of Fe₃O₄ nanoparticles much better than uncoated nanoparticles alone and leads to higher magnetocrystalline anisotropy. A study by Liu et al. [142] suggested the possibility of increasing SAR by decreasing the surface-coating thickness using highly monodispersed Fe₃O₄ nanoparticles with different polyethylene glycol-coating thickness. The increase in SAR was explained as due to a decrease in coating thickness leading to an increased Brownian loss, improved thermal conductivity, as well as improved dispersion. It should be noted that the heating performance of the nanoparticles depends on the medium as well. Whenever nanoparticles encounter biological systems, interactions take place between their surfaces and biological components such as proteins, membranes, phospholipids, and DNA forming the so-called protein corona around the nanoparticles [143, 144]. The formation of corona depends on the surface properties of the particles [145, 146] and can influence the aggregation behavior of nanoparticles in biological media, which in turn can affect their performance for desired applications. Therefore, apart from the optimization of the properties of the magnetic core and surface coating for high-performance MHT, it is necessary to ensure its performance in the physiological environments.

In the work reported by Khandhar et al. [147], authors use poly(maleic anhydride-alt-1-octadecene)-poly(ethylene glycol) (PMAO-PEG), an amphiphilic polymer-coated Fe₃O₄ nanoparticles of three different sizes, 13, 14, and 16 nm, to study the MHT efficiency in cell growth medium (CGM) similar to biological environment. Their results showed an increase in hydrodynamic sizes in all three samples upon exposure to CGM. SAR reduced (30%) only in 16-nm size sample, while other two samples did not exhibit any significant decrease in SAR. The authors suggest that the increase in hydrodynamic volume prolongs Brownian relaxation while Néel relaxation is unaffected. Hence, in 13- and 14-nm samples where SAR is mainly due to Néel relaxation, SAR was not affected. But in the 16-nm samples, in which there is a contribution from Brownian relaxation to heat dissipation, the SAR dropped due to the increase in Brownian relaxation. We have investigated the magnetic hyperthermia efficiency of dextran and citric acid (CA)-coated Fe₃O₄ ferrofluids in cell growth medium, which contains
serum proteins similar to physiological environments. From the stock solutions (25 mg/ml), 3 mg/ml concentration of dextran and citric acid-coated ferrofluids samples were prepared using CGM and deionized water (DI) water. The ferrofluid samples were subjected to an AC field of 235-Oe amplitude at the frequency of 375 kHz. The SAR of dextran-coated samples in DI and CGM was estimated to be 63 and 72 W/g, which indicates that their performance is not much affected by the medium if not enhanced. However, SAR values obtained for CA-coated samples in DI and CGM, 78 and 38 W/g, implies that their efficiency is heavily reduced when exposed to physiological environments.

7. Gd-doped SPION as a potential theranostic agent

The multifunctionality of SPION makes them a good candidate for theranostics. One such approach to integrate diagnostic imaging and therapeutic function is to develop SPION as an MRI/drug delivery platform. Yu et al. [148] reported that PEG-coated iron oxide nanoparticles when loaded with Dox provide a therapeutic capability. Following their injection into a mouse, Dox-modified magnetic nanoparticles accumulate in the tumor and the nanoparticles were imaged by using $T_2$ MRI. The contrast associated with the tumor changes from light to dark at 4.5 h post injection and the growth rate of the tumor mass was decreased in the nanoparticle-injected mice compared to that of a control group. In another work, Lee et al. [149] developed PEG-stabilized Fe$_3$O$_4$ nanocrystals on dye-doped mesoporous silica nanoparticles and Dox was loaded into the pores. Here, SPIONs work as a contrast agent in MRI, the dye molecule imparts optical imaging modality, and Dox induces cell death. In a similar approach, Kim et al. [150] developed a core-shell structure consisting of single Fe$_3$O$_4$ core and mesoporous silica shell for MR and fluorescence imaging which also has the potential to be used as a drug carrier. Hayashi et al. [92] reports a study done on SPION conjugated with folic acid as targeting ligand and Tamoxifen as anticancer drug. The drug release was triggered by heat generated by SPION in an AC magnetic field, hence performing drug delivery and hyperthermia simultaneously.

The incorporation of MHT and imaging modalities [151] has been investigated widely as well. One such study is reported by Hayashi et al. [152] in which authors have investigated SPION for cancer theranostics by combining MRI and magnetic hyperthermia through a set of in vivo experiments. They show that FA- and PEG-modified SPION nanoclusters accumulated locally in cancer tissues within the tumor and enhanced the MRI contrast. Furthermore, they report that with MHT, the tumor volume of treated mice was reduced to one-tenth that of the control mice. Also, Gd-doped Fe$_3$O$_4$ nanoparticles have the potential to act as an effective MHT agent [153] in addition to their use as a $T_1$-$T_2$ dual-modal contrast agent for MR imaging. Gd(III) is known to oppose net magnetic moment of Fe(III)/Fe(II) in oxides, reducing magnetization [154–156]. Therefore, Gd doping may reduce the hyperthermia efficiency, but by using the correct amount of doping, one can always explore the possibility of using them as both MRI contrast agents and hyperthermia mediators. However, there are very few studies done on the MHT efficiency of Gd-doped Fe$_3$O$_4$ nanoparticles [157, 158]. Both the studies report higher SAR values for Gd-doped Fe$_3$O$_4$ nanoparticles compared to the reported values for undoped samples. In our work presented
here, we have investigated the MHT efficiency of Gd$_{0.075}$Fe$_{2.925}$O$_4$ nanoparticles for possible use as a theranostic agent.

### 7.1. Synthesis and characterization of Gd-doped SPION

Gd$_{0.075}$Fe$_{2.925}$O$_4$ nanoparticles were synthesized by coprecipitation method. For a typical synthesis of Gd$_{0.075}$Fe$_{2.925}$O$_4$, aqueous solution of FeCl$_2$.4H$_2$O, FeCl$_3$.6H$_2$O and Gd(NO$_3$)$_3$ was mixed in a molar ratio of 1.00:1.925:0.075 in 25-ml volume followed by the addition of 250 ml of 1 M NH$_4$OH. The synthesized nanoparticles were then coated with dextran according to the method outlined by Arachchige et al. [140]. From the structural investigation, it was observed that Gd doping does not alter the Fe$_3$O$_4$ crystal structure significantly (Figure 4). Using several intense X-ray diffraction (XRD) peaks and the Debye-Scherer equation, the crystallite sizes of the Fe$_3$O$_4$ and Gd$_{0.075}$Fe$_{2.925}$O$_4$ nanoparticle samples were determined to be 11.7 ± 0.6 and 14.9 ± 0.5 nm, respectively. This increase in the crystallite size is consistent with the previous studies on Gd doping in spinel structures [159].

TEM images of the two samples are shown in Figure 5. The undoped sample consists of roughly spherical nanoparticles with smaller polydispersity, whereas the Gd-doped sample exhibits nanoparticles with rough edges with wider size distribution.

The magnetic properties of the synthesized powder as well as the ferrofluid samples are determined by analyzing the $M(H)$ curve. The M-H data for undoped and Gd-doped Fe$_3$O$_4$ ferrofluid samples, recorded at room temperature, are shown in Figure 6. The sigmoidal shape of the $M(H)$ curves with nearly zero hysteresis confirms the superparamagnetic nature of these nanoparticles at room temperature. The saturation magnetization of Fe$_3$O$_4$ nanoparticles is measured to be ~72 emu/g, whereas that of Gd-doped Fe$_3$O$_4$ nanoparticles is reduced to ~52 emu/g. This reduction in saturation magnetization at room temperature agrees with the observations in other reported studies [123, 158] and can be attributed to the fact that magnetic Fe$^{3+}$ ions get replaced by the Gd$^{3+}$ ions in the octahedral sites of the inverse spinel structure.

![Figure 4. X-ray diffraction patterns of as-prepared Fe$_3$O$_4$ and Gd$_{0.075}$Fe$_{2.925}$O$_4$ nanoparticles [160].](image-url)
It is observed that the doping of Gd$^{3+}$ ions into Fe$_3$O$_4$ spinel has significantly influenced the average crystallite size and the saturation magnetization. The $M(H)$ curve for an ensemble of noninteracting superparamagnetic nanoparticles described by a log-normal distribution function, $f(D)$, can be fitted using the following expression:

$$M(H) = M_s \int_0^{\infty} f(D) V L(x) dD$$

where $L(x) = \coth \frac{x}{2}$ is the Langevin function, $x = (M_s VH)/k_B T$, $M_s$ is the saturation magnetization, and $V$ is the volume of the particle. The fitted particle size was inconsistent with the

Figure 5. TEM images of (a) Fe$_3$O$_4$ and (b) Gd-Fe$_3$O$_4$ nanoparticles [160].

Figure 6. $M$ versus $H$ curves for two ferrofluid samples fitted with Eq. (5). The inset shows the resulting particle size distribution obtained for the two samples [160].
observed XRD data, and it was necessary to introduce the magnetic dipolar interaction effects through a phenomenological temperature, $T^*$, as described in our recent work [140]. The best-fit parameters for the two samples are shown in Table 2, and Figure 6 shows the fitted data. The inset in Figure 6 shows the magnetic core size distributions for two ferrofluid samples. The fitting of $M(H)$ data with Eq. (5) clearly shows that Gd-doped Fe$_3$O$_4$ nanoparticles have a higher average magnetic core size with a larger size distribution (14.6 ± 3.7 nm) and lower saturation magnetization (52 emu/g) compared to the undoped Fe$_3$O$_4$ nanoparticles (11.7 ± 1.9 nm, 72 emu/g). Both the ferrofluid samples exhibit similar strength of magnetic dipolar interaction ($T^* \approx 80–100$ K).

MHT measurements were carried out on the dextran-coated Gd-doped as well as undoped Fe$_3$O$_4$ ferrofluid samples at a field of 235 Oe and at a frequency of 375 kHz. The heating curves for two samples are shown in Figure 7(a), and from the plot it can be observed that the initial heating rates for the two samples are approximately the same. From these heating curves, the SAR values were obtained as a function of temperature taking into account heat loss as described elsewhere [134]. Figure 7(b) presents the corrected experimental SAR data as a function of temperature for both the undoped and Gd-doped samples. Within the experimental error, the room temperature SAR values for Gd-doped Fe$_3$O$_4$ and undoped

<table>
<thead>
<tr>
<th>Ferrofluid sample</th>
<th>$M_s$ (emu/g)</th>
<th>$D_0$ (nm)</th>
<th>$\sigma$</th>
<th>$T^*$ (K)</th>
<th>$D_{avg}$ (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe$_3$O$_4$</td>
<td>72</td>
<td>11.6</td>
<td>0.15</td>
<td>80</td>
<td>11.7 ± 1.9</td>
</tr>
<tr>
<td>Gd -Fe$_3$O$_4$</td>
<td>52</td>
<td>14.2</td>
<td>0.23</td>
<td>100</td>
<td>14.6 ± 3.7</td>
</tr>
</tbody>
</table>

Table 2. Fitting parameters obtained from the $M(H)$ fitting with modified Langevin function using $T^*$.

Figure 7. (a) Heating profiles of Fe$_3$O$_4$ and Gd-doped Fe$_3$O$_4$ ferrofluid samples under an AC magnetic field amplitude of 235 Oe and at a frequency of 375 kHz. (b) The temperature dependence of net SAR for two ferrofluid samples. The black line shows the theoretical fitting of the experimental data with the linear response theory [160].
Fe₃O₄ ferrofluid are very similar. The temperature-dependent SAR values were fitted to the linear response theory incorporated with the interactions and size distribution [140]. The solid lines in Figure 7(b) are the best fits to the experimental SAR data, using the particle size distribution parameters and $T^*$ values given in Table 2 and treating the anisotropy constant, $K$, as a fitting parameter. The SAR fitting yields a somewhat smaller anisotropy constant ($\sim 12$ kJ/m$^3$) for Gd-doped ferrofluid compared to that of undoped ferrofluid ($\sim 21$kJ/m$^3$). It is interesting to note that both samples have similar SAR values in spite of a smaller anisotropy constant and the saturation magnetization for the Gd$_{0.075}$Fe$_{2.925}$O$_4$ sample compared to the undoped sample. The expected lowering of SAR in Gd$_{0.075}$Fe$_{2.925}$O$_4$ is probably offset by a larger particle size in this sample, as the SAR would increase with increasing particle size up to a critical size [140]. By fine-tuning the composition of Gd-doped Fe$_3$O$_4$ nanoparticles, we may achieve a higher SAR value.

In summary, the Gd doping on the Fe$_3$O$_4$ nanoparticles affects the morphology and the magnetic properties of Fe$_3$O$_4$ nanoparticles considerably but the magnetic hyperthermia efficiency of the samples was about the same within the experimental uncertainties. The possibility of using Gd-doped Fe$_3$O$_4$ nanoparticles as a dual-modal $T_1$-$T_2$ contrast agent is being currently explored by others and our magnetic hyperthermia results demonstrate that this material is a potential candidate for multimodal contrast imaging and cancer treatment by hyperthermia. However, further research is necessary to optimize the amount of Gd doping to enhance SAR for cancer treatment and to be used as a theranostic agent.

8. Conclusions

In this chapter, we have discussed various approaches to exploit the multifunctionality of SPION for cancer theranostics. We have given a brief background on the nanoparticle magnetism, followed by a description of commonly used synthesis methods and surface-functionalizing strategies. Three major applications of Fe$_3$O$_4$ nanoparticles in drug delivery, MRI, and MHT, including our recent work on Gd-doped SPION as a possible theranostic agent, are described. This chapter also addresses the recent work on integrating the individual diagnostic and therapeutic approaches to develop SPION-based theranostic platform.

Despite the exciting progress, SPION is far from meeting clinical standards as theranostic agent. It has its own promises and advantages, but there are still some disadvantages to be overcome. These include target specificity as drug carriers, optimizing the heating efficiency and aim for sufficient heating using minimum dosage, preventing the overheating in MHT, and issues of biocirculation, biodistribution, and bioelimination within the biological system. In summary, although in theory, SPION is a perfect vehicle in the development of theranostic nanomedicine, more research is required to overcome its disadvantages and this should be the main focus of the next stages of investigation.
9. Future directions

In recent years, the research in the field of theranostics has brought many diverse fields together for targeting, imaging, and therapy for a deadly disease like cancer. These fields include physics of magnetism, chemistry of synthesis, material science of structure-property relationship, surface science for functionalization, biomedical engineering in MRI and radiofrequency activation and treatment, and biology for understanding the behavior of cancer cells. SPIONs have play ed a key role in this application as visual, imaging, and therapeutic agent. Several studies have shown promising results; however, many challenges still remain in moving theranostic applications from laboratory settings to clinics. Two major challenges we face are low efficacy and toxicity of SPION. For in vivo applications, the amount of SPION used (several hundred microgram/ml) usually produces undesirable toxic side effects. The smaller concentration, on the other hand, is not sufficient for imagining and therapeutic action of the material. It is well known that the size, shape, and surface modifications influence the performance of SPION. There is a lack of information about the combined effects of these parameters in the clinical applications. Also, we do not have a clear understanding of controlling the delivery of SPION to a specific target in the body by using external magnetic field gradient. It has been found that some particles would end up accumulating in other parts of the body such as liver, spleen, kidney, and lungs along with the specific intended location. We do not know how they will affect those nonspecific organs and how long they will stay there. In magnetic hyperthermia therapy, measuring precise temperature at the tumor site and adjusting particle properties with frequencies and amplitude of the external field for apoptosis/necrosis of cancer cells without affecting normal tissues are challenges that researchers and clinicians face every day. There have been promising results in treating prostate and skin cancers with magnetic fluid hyperthermia but no real efforts have been made to treat deep tumors such a pancreas and liver with SPION. In order to make progress with these therapies, research is needed in the development of new materials that have higher reflexivity, better thermal activation properties, and have better coating materials to improve the bio-distribution and biocompatibility for in vivo applications. Most imperatively, we need data on large animal studies before theranostics can make a fruitful transition from research laboratories to the clinics, and so on.

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