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REVIEW

In vivo Biodistribution and Clearance of Magnetic Iron Oxide Nanoparticles for Medical **Applications**

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Abstract: Magnetic iron oxide nanoparticles (magnetite and maghemite) are intensively studied due to their broad potential applications in medical and biological sciences. Their unique properties, such as nanometric size, large specific surface area, and superparamagnetism, allow them to be used in targeted drug delivery and internal radiotherapy by targeting an external magnetic field. In addition, they are successfully used in magnetic resonance imaging (MRI), hyperthermia, and radiolabelling. The appropriate design of nanoparticles allows them to be delivered to the desired tissues and organs. The desired biodistribution of nanoparticles, eg, cancerous tumors, is increased using an external magnetic field. Thus, knowledge of the biodistribution of these nanoparticles is essential for medical applications. It allows for determining whether nanoparticles are captured by the desired organs or accumulated in other tissues, which may lead to potential toxicity. This review article presents the main organs where nanoparticles accumulate. The sites of their first uptake are usually the liver, spleen, and lymph nodes, but with the appropriate design of nanoparticles, they can also be accumulated in organs such as the lungs, heart, or brain. In addition, the review describes the factors affecting the biodistribution of nanoparticles, including their size, shape, surface charge, coating molecules, and route of administration. Modern techniques for determining nanoparticle accumulation sites and concentration in isolated tissues or the body in vivo are also presented. Keywords: magnetic nanoparticles, iron oxide nanoparticles, biodistribution, biodegradation

Introduction

Magnetic nanoparticles/iron oxide nanoparticles (MNPs/IONPs) (magnetite Fe₃O₄ and maghemite γ -Fe₂O₃) have been under intensive investigation since they show great potential for biomedical applications, such as tumors imaging (MRI),¹⁻³ radiolabelling and internal radiotherapy,⁴ hyperthermia,^{5,6} gene therapy,⁷ biomolecules separation^{8,9} and drug delivery.^{9,10} Drug delivery is a vast field of MNPs applications as they possess a relatively large surface that enables binding and carrying other compounds, which can be directed using an external magnetic field to specific places in the body.¹¹ Additionally, nanoparticles depending on their size, can penetrate almost all tissues. Importantly, drugs immobilized on MNPs usually gain more beneficial properties, such as better solubility in water¹² and higher cytotoxicity to cancer cells.^{13,14}

The uniqueness of iron oxide nanoparticles lies in their ferromagnetic nature. They consist of areas of permanent magnetization (so-called magnetic domains), generating a magnetic field around them (like small magnets). Ferromagnets are ferromagnetic below the Curie temperature and paramagnetic above. Magnetization is changing the direction of magnetization in domains or the boundaries of domains. All elementary magnetic dipoles align with an external magnetic field at magnetic saturation. The external magnetic field can guide magnetite and maghemite nanoparticles to specific locations in the body.^{15,16} In the case of magnetic-targeted drugs, such as anti-cancer drugs, this results in lower toxicity and lower doses. Ferromagnetism distinguishes MNPs from other types of nanoparticles used in medicine, such as gold (AuNPs), silver (AgNPs) nanoparticles, or polymer nanoparticles (PNPs).

Graphical Abstract



On the surface of AuNPs, electrons oscillate under the influence of light (surface plasmon resonance). AuNPs can scatter light intensely and are photostable, which makes these particles attractive imaging tools.¹⁷ They are also used as contrast agents in computed tomography.¹⁸ Among nanoparticles, AuNPs, and MNPs have also gained prominence as nanoradiopharmaceuticals for cancer imaging and therapy due to their biocompatibility, functionalization capabilities, and radiolabeling potential.^{19–25} These targeted nanobrachytherapy approaches offer less invasiveness and improved radiation dose distribution compared to conventional treatments. Silver nanoparticles also show radiosensitizing properties.²⁶ Unfortunately, the main disadvantage of metallic nanoparticles is their non-biodegradability and toxicity. For example, the mice given 8 mg/kg AuNPs per body developed fatigue and a crooked spine, most of which died within three weeks.²⁷ In addition, Cho et al studied that PEGylated AuNPs (maximum dose 4.26 mg/kg) caused apoptosis of liver tissue.²⁸ An interesting type of nanoparticle in medical applications is also polymer nanoparticles (PNPs), such as nanospheres with a matrix structure, nanocapsules, dendrimers, and nanomicelles.^{29–31} Some of them, similarly to

magnetic nanoparticles, enable drug release in a controlled and sustained manner and increase the solubility of lipophilic drugs.³¹ Unfortunately, most PNPs have not been adequately tested regarding toxicology. Furthermore, there are difficulties in increasing the scale of their production, and the costs of producing some of them are very high.³¹

Magnetic iron oxide nanoparticles, unlike other metallic particles, are mainly biodegradable. The biodegradation mechanism of MNPs is believed to be analogous to ferritin metabolism, which is digested by lysosomal enzymes to release iron ions.³² Many studies indicate that magnetic nanoparticles show low toxicity, causing only short-term, temporary changes, eg, oxidative stress, but do not contribute to permanent damage to organs.^{33–40} Only extremely high doses (500 mg Fe/kg) caused pathological changes in the liver and spleen.⁴¹

Biodistribution of magnetic nanoparticles in the organism is an important parameter, the knowledge of which makes it possible to reduce the toxicity of nanostructures, which is a result of, inter alia, their undesirable distribution, as well as to increase the expected functionality of MNPs in a selected, targeted organ or tissues.⁴² The parameter determines the amount and place of MNPs accumulation in various body organs during or after their application in vivo.⁴³ Biodistribution is closely related to the physicochemical properties of nanoparticles, such as size, coating molecules, and surface charge.^{43,44}

The commonly used approach for administering MNPs is an intravenous injection, especially for their use as MRI contrast agents and in the case of clinical oncology.⁴³ The intravenously administered nanoparticles are selectively taken up by the liver and spleen, constituting the main pathway for removing MNPs from the body. Both organs form the mononuclear phagocytic system (MPS), also known as the reticuloendothelial system (RES). It should be noted that although MPS and RES nomenclatures are most often used interchangeably concerning macrophages, RES comprises only sinusoidal epithelial cells in the liver.⁴⁵ MPS consists of circulating monocytes in the blood and macrophages in various organs such as the liver, spleen, lymph nodes, bone marrow, lungs, and brain.⁴⁶ Among the specialized macrophages, there are, among others, Kupffer cells present in the sinusoidal capillaries of the liver, microglial cells found in the brain, as well as macrophages inhabiting secondary lymphatic organs, such as the spleen, and more specifically the red pulp and marginal zones of this organ.^{46,47} By phagocytosis and engaging other macrophages from the bloodstream, macrophages cleanse the body of pathogens, damaged and old cells, and foreign bodies, including MNPs.^{47,48} Although the liver and spleen are the main sites of nanoparticle uptake,⁴⁹ when injecting high doses of MNPs, their excess is accumulated in other tissues and organs such as lungs, heart, or adipose tissue.⁵⁰ It should be emphasized that the opsonization of nanoparticles precedes the uptake in the liver and spleen.^{51,52} The opsonization process involves attaching specific proteins in the bloodstream to the surface of MNPs. These marking proteins are primarily immunoglobulins (IgA and IgM), complement components (C3, C4, C5),⁵³ and blood serum proteins (albumins, fibrinogen, fibronectin, C-reactive protein, type-I collagen).⁵⁴ Due to the protein labeling, the nanoparticles are recognized by macrophages.

The Main Organs and Tissues Involved in the Accumulation of Magnetic Nanoparticles

Liver

Blood from both the portal vein and hepatic artery mixes in the hepatic sinusoids. Sinusoidal capillaries, which are highly fenestrated with an average pore size of 100–180 nm, contain specialized macrophages called Kupffer cells (Figure 1).⁵⁵ Kupffer cells comprise the largest population of tissue macrophages in the organism and play a crucial role in the immune response efficiently phagocytizing pathogens entering from the bloodstream, including MNPs. Hepatocytes are separated from the sinusoids by a region called Disse space.⁵⁶ Ito cells in Disse space store fat-soluble vitamins. Kupffer cells quickly uptake large nanoparticles (> 100 nm) or agglomerates,⁵⁷ whereas smaller nanoparticles (< 100 nm) are captured in the Disse space. Hence, if they are provided with a suitable "shell", they can accumulate in hepatocytes.⁵⁸ Distribution in hepatocytes can be increased inter alia by attaching to nanoparticles of molecules with high affinity for these cells, such as linoleic acid⁵⁹ or by functionalization of the MNPs surface using antifouling molecules such as PEG⁶⁰ which reduce the degree of opsonization.



Figure I In the liver, Kupffer cells can quickly uptake large nanoparticles (> 100 nm), whereas smaller nanoparticles (< 100 nm) are captured in the Disse space from where, if suitably functionalized, they can also accumulate in hepatocytes.

Many reports in the literature indicate that the liver is the primary capture and biodistribution site for magnetic nanoparticles, both in the short and long term. A series of detailed studies in this field have confirmed it. For example, Tate et al⁶¹ tested the biodistribution of two types of magnetite nanoparticles coated with hydroxyl-ethyl starch with an average hydrodynamic diameter of 110 nm and 20 nm in mice models. The results showed that the liver next to the spleen was the primary site of iron accumulation after 72 h for larger-sized nanoparticles (approx. 1.6 mg Fe/g tissue) (Figure 2A). Their level in this organ was still high even after 14 days after injection (Figure 2B). In contrast, image quantification of the heart, brain, and lung showed no significant difference in mean percent iron content.

Other studies have shown that polyacrylic-coated nanoparticles with an average diameter of approx. 10 nm after intravenous injection in mice models were also mainly accumulated in the liver.³³ The iron level in this organ 24 h after injection was about 2.5 mg/g of organ at a dose of 50 mg/kg body weight. This value was equivalent to approx. 6.5% liver weight/per total weight ratio, whereas in the case of the spleen, the ratio was around 0.41% spleen weight per total weight ratio. The high biodistribution of magnetic nanoparticles was also confirmed in the rabbit liver.⁶² The authors used iron oxide nanoparticles covered with phosphated starch with a hydrodynamic diameter of 100–250 nm, which were intraarterially or intravenously injected. Shortly after application, the amount of the detected nanoparticles in the liver was 44% of the whole amount of the MNPs administered in the case of intraarterial injection and 73% in the case of intravenous injection.

Moreover, these values were not affected by the sizes of nanoparticles. Another study investigated the biodistribution of ultra-small superparamagnetic iron oxide nanoparticles used for MRI in BALB mice.⁶³ The nanoparticles were labeled with radioisotope ^{99m}Tc and administered via intravenous injection (the average diameter of ^{99m}Tc-MNPs was 41 nm). About 75% of injected doses were found in the spleen and liver at 15 min post-injection. While about 24% injected dose remained in the liver after 48 h post-injection, it was just about 3% in the spleen. Wang et al investigated the biodistribution of ¹¹¹In-radiolabeled PEGylated superparamagnetic iron oxide nanoparticles with a size range of 5–25 nm in the mouse.⁶⁴ The results showed that the liver exhibited the highest radioactivity concentrations (10 min, 30 min, 1 h, 2 h, and 24 h), followed by the spleen. After 24 h, radioactivity reached the value of 37% per gram of organ (for comparison – the radioactivity values after that time in the spleen and kidney were 21% and 8%, respectively). In summary, presented studies show that the liver is the initial filtration point where magnetic nanoparticles accumulate.



Figure 2 ICP-MS liver iron content after 72 hours (A) and after 14 days (concerns 110 nm-sized MNPs) (B). Percent of iron-stained pixels in liver tissue (iron-stained pixels were identified by calculating the Euclidian distance of each pixel to an average iron-stained RGB value given from a Prussian Blue iron-stained control slide). Used with permission of SPLE, from In Vivo Biodistribution of Iron Oxide Nanoparticles: An Overview. Energy Based Treat Tissue Assess, Tate JA, Petryk AA, Giustini AJ, Hoopes PJ. 2011; permission conveyed through Copyright Clearance, Inc.⁶¹

Because of the high accumulation in the liver, iron oxide nanoparticles have been successfully used as MRI T_2 contrast agents in this organ.^{65,66} In particular, it is beneficial for detecting potential liver tumors as MNPs are taken up by Kupffer cells in healthy parts of the liver generating dark contrast in T_2 -weighted MRI. In contrast, cancer cells not containing Kupffer cells appear as bright regions.⁶⁷ Xu et al have shown that glutathione-responsive hyaluronic acid-coated magnetic nanoparticles (H-MNPs) can be used for highly sensitive liver metastases (LMs) diagnosis.⁶⁸ The detection of small LMs is still challenging because of the subtle differences between normal liver tissue and small metastases. The authors demonstrated that H-MNPs greatly enhanced the signal of MRI in tumor metastases as a T_1 contrast agent. In contrast, they substantially decreased the signal of the liver as a T_2 contrast agent as they aggregated into clusters upon the high glutathione level in the liver.

Consequently, MRI contrasted by H-MNPs clearly distinguished metastatic tumors (bright) from surrounding liver tissues (dark). Moreover, there are reports that magnetic nanoparticles can be applied for liver fibrosis theranostics.⁶⁹ For example, Saraswathy et al tested citrate-coated magnetic nanoparticles with an average diameter of 12 nm and observed

an apparent dose-dependent contrast enhancement in a T_2 -weighted in vivo MR imaging of liver fibrosis model in rodents.⁷⁰ On the other hand, due to such a significant capture of nanoparticles by the liver, their delivery to other tissues and target organs may be insufficient.⁷¹ Thus, as there is the possibility of synthesizing nanoparticles of different sizes, morphology, and surface chemistry, there is a need for more extensive research into the type of interactions of magnetic nanoparticles with the liver to design nanoparticles according to therapeutic goals.

Lymphoid Organs: Spleen and Lymph Nodes

The spleen and lymph nodes are secondary lymphatic organs that function as biological filters protecting the body against microbes and cancerous cells.⁷² They are also the site of the proliferation and maturation of T and B lymphocytes. The outer layer of the spleen is a compact connective tissue capsule, while its parenchyma consists of two types of interpenetrating tissues called white and red pulp (Figure 3A). Blood through the splenic artery first enters the white pulp and then, through smaller arterioles, reaches the red pulp, which constitutes 75% of the volume of the entire organ.⁷³ The spleen's primary immunological function is to remove pathogens and foreign particles (including MNPs) and to destroy old and senescent red blood cells (RBC) digested by phagocytic macrophages in the red pulp. The iron from RBC is removed and transported to be stored as ferritin.

Macrophage phagocytic pathogens, including magnetic nanoparticles, are found in the marginal zone of the spleen, which separates the red and white pulp. There is also a population of macrophages in the white pulp, performing functions similar to those in lymph nodes.⁷⁴ In red pulp, macrophages play the role of cleansing the blood of residual erythrocytes and are involved in iron recycling. On the other hand, there are reports that magnetic nanoparticles, although mainly phagocytosed by macrophages in the marginal zone, are also captured by macrophages in the red pulp of the spleen.^{75,76}

Lymph nodes vary in size and shape and are found clustered at specific locations throughout the body. Each lymph node is divided into several regions: fibrous capsule, outer (nodular) cortex rich in B-cells, inner cortex (paracortex) rich in T-cells, and medulla – the central inner part of the node that contains large numbers of fixed phagocytic macrophages (Figure 3B).⁷² MNPs can get into the lymph surrounding the tissues and then, through the lymphatic vessels, are directed to the lymph nodes, where they are trapped in a sinusoidal mesh containing macrophages.^{77–79} The significant fraction of



Figure 3 Schematic structure of spleen (A) and lymph node (B).

the intravenously injected MNPs is usually first captured by the liver and spleen. However, regional lymph nodes may be the initial filter point if magnetic nanoparticles are administered intramuscularly or subcutaneously.

A series of reports demonstrate a high accumulation of magnetic nanoparticles in the spleen. For example, about 10 nm-sized curcumin-coated MNPs administered intravenously in repeated doses in mice model reached a high level in the spleen 3 days after the final injection (about 110 μ g Fe/dl; for comparison - this value in the case of the liver was about 60 μ g Fe/dl).³⁴ In other studies, oleic acid-pluronic-coated MNPs were administered intravenously in rats. The Fe level was the highest in the spleen, reaching approximately 1200 μ g/g of tissue after 21 days of injection.³⁵ The accumulation of these nanoparticles in the spleen was the highest compared to the other organs tested (Figure 4).

Kidneys and Non-Phagocytizing Clearance Pathway

The kidney, a bean-shaped organ, has a hilum, the passageway for the renal vessels and the ureter. The primary function of this organ is the filtration of fluids from the bloodstream, discarding the excess water and ions and eliminating nitrogenous waste, toxins, and drugs from the body. Each kidney contains over a million tiny structures called nephrons responsible for forming urine. Nephrons contain glomerular vessels that involve infiltration.

The intravenously injected MNPs enter the kidney with an artery in the renal hilum and reach glomeruli in nephrons. In glomeruli, cells called podocytes wrap around capillaries. If MNPs can get through the glomerular endothelium and



Figure 4 Changes in iron levels in different tissues following intravenous injection of MNPs. The injected dose of iron was 10 mg/kg in 100 μ L of saline. Data presented as mean ± sem (n = 4). *p < 0.05 or **p < 0.005 vs saline. Reprinted with permission from Jain TK, Reddy MK, Morales MA, Leslie-Pelecky DL, Biodistribution, Clearance, and biocompatibility of iron oxide magnetic nanoparticles in rats. Mol Pharm. 2008;5(2):316–327. Copyright©2008, American Chemical Society.³⁵

fenestrations between the podocytes, they are eventually extracted in urine via the ureter. This applies to nanoparticles smaller than 10-15 nm.^{44,80}

Many studies have been performed showing renal clearance of small magnetic nanoparticles. For example, Kim et al developed iron oxide nanoparticle-based T_I musculoskeletal contrast agents having an inverted core–shell structure of a dextran core and an iron oxide shell (i-MNPs) and a size of 3 nm.⁸¹ One of the beneficial features of these nanoparticles is their compact hydrodynamic diameter, which is smaller than the renal filtration size limit. Intra-articularly injected i-MNPs were entirely excreted through renal clearance (Figure 5). According to the authors, it was the first successful demonstration of the renal clearance of intra-articularly injected nanomaterials.

Other studies showed the use of MNPs for the thermoablation of malignant kidney tumors in rabbits.⁸² The nanoparticles' core size was 9–12 nm. In most cases, the applied ferrofluid was concentrated within the tumor mass. Additionally, a small portion of the ferrofluid was found in the renal pelvis or collecting duct. However, one should be careful when interpreting the results of the biodistribution of nanoparticles in the kidney, especially if that is the case with MNPs labeled with fluorophores. It is possible to locate the particles of the fluorophores themselves released from the nanoparticles as within the GBM (glomerular basement membrane), the nanoparticles can be disassembled by the numerous heparan sulfate molecules (Figure 6).⁸³ Once disassembled, the nanoparticle components can cross the remainder of the GBM and the podocyte filtration slits and enter the urinary space.

Furthermore, the fluorescent signal in the kidneys or MRI contrast enhancement in this area does not necessarily mean the uptake of MNPs by the glomeruli but only the result of the circulation of nanoparticles in the capillaries and arterioles of the renal cortex. The results obtained for PAMAM-dendrimer-coated MNPs with a core size of 10 nm and a huge hydrodynamic size of 242 nm may confirm the above.⁸⁴ The studies' authors demonstrated that kidneys were the



Figure 5 Renal excretion of intra-articular injected i–IONPs (i-MNPs): cumulative urinary iron content of male rats after intra-articular injection of i–IONPs (i-MNPs) and saline (inset: representative photos of urine samples of saline-injected (left) and i–IONPs-injected (right) rats. n = 5 biologically independent rats for each group. Error bars represent the standard deviation (***P < 0.001). Reprinted with permission from Wook KJ, Cheong J, Cheong H, et al. Iron oxide-coated dextran nanoparticles with efficient renal clearance for musculoskeletal magnetic resonance imaging. ACS Appl Nano Mater. 2021;4(12):12943–12948. Copyright©2021, American Chemical Society.⁸¹



Figure 6 Extraction pathway of the MNPs or their degradation products through the kidney. MNPs smaller than 10–15 nm, the released coating molecules, degradation byproducts, and therapeutic agents can cross the GMB and pores between podocytes and enter the urinary space.

main distribution sites of such nanoparticles. However, it is supposed that it could have been, for example, the result of the loss of coating molecules by magnetite 10 nm-sized cores.

Lungs

The respiratory part of the respiratory system includes bronchial tubes, alveolar ducts, and pulmonary alveoli.⁸⁵ During respiration, oxygen enters the lungs by diffusion through the capillaries surrounding each alveolar sac. Pulmonary macrophages are present in the alveoli. They can be observed in the interalveolar septa (interstitial macrophages) and inside and on the surface of the pulmonary epithelium (alveolar macrophages) surface. These cells phagocytose foreign bodies and induce an immune response. Alveolar macrophages often leave the alveoli and travel to the bronchioles, moving towards the upper respiratory tract, while others enter the lymphatic system.

Until recently, it was mistakenly believed that MNPs are poorly recognized by the body's defenses and thus ineffectively removed from the lungs due to the small accumulation of phagocytes.⁸⁶ It has been demonstrated that pulmonary phagocytosis responds very effectively to metallic nanoparticles' deposition, including magnetic nanoparticles.⁸⁷ Nanometric MNPs are removed from the lung tissue faster than micrometric particles, probably due to more active phagocytosis of nanostructures. This is of great importance because nanoparticles administered by inhalation are deposited mainly in the lungs.

A series of studies concern the biodistribution MNPs in organs, including lungs. For example, Edge et al investigated the biodistribution of 12 nm- and 15-sized superparamagnetic iron oxide nanoparticles in pigs.⁸⁸ Using pEPR technique analysis, they observed a dose-dependent accumulation for both sizes of nanoparticles in pig lungs compared to controls after 5 hours after injection. This was confirmed by histological staining with Prussian blue, showing MNPs localized in alveolar tissue. Other studies on the biodistribution of hydroxyl-ethyl starch-coated MNPs in mice using ICP-MS have shown the presence of nanoparticles in lungs after 4 hours after intravenous injection (about 0.15 mg Fe/g tissue). However, the number of deposited MNPs decreased over time, reaching a value of about 0.1 mg Fe/g tissue after 72 h.⁶¹ After 14 days, the nanoparticles in the lungs were undetectable This confirms the thesis that MNPs in these organs are phagocytosed rapidly, thus reducing the risk of their side effects.

Research is also being conducted on using magnetic nanoparticles as drug carriers in the lungs. For example, the heavily supplied with capillaries lung tissue was exploited to deliver positive charge-coated MNPs containing the anticancer drug irinotecan.⁸⁹ The endothelium is lined richly with negatively charged heparin sulfate proteoglycans⁹⁰ to bind the positively charged growth factors and proteins and positively charged nanoparticles, for example, coated with positively charged lysine. Magnetic nanoparticles were also studied as potential early diagnosis lung cancer agents.⁹¹ Moreover, Tan et al proposed using magnetic nanoparticles by inhalation as drug carriers directly to the primary target site.⁹² This allows for avoiding the risks of systemic toxicity.

Heart

Tissue-resident macrophages have also been observed in the heart. Cardiac macrophages are self-proliferating and independent of blood monocyte-derived macrophages.⁹³ The macrophages in the heart are very heterogeneous and show phenotypic and functional adaptation depending on the surrounding environment and aging. Two main groups can be distinguished, including classical "inflammatory" macrophages with pro-inflammatory properties (M1) and alternatively "anti-inflammatory" macrophages (M2) associated with cell proliferation and tissue repair. The essential functions of M1 macrophages are phagocytosis and destruction of bacteria, elimination of neoplastic cells, and production of pro-inflammatory cytokines.⁷⁴ However, the exact role of cardiac macrophages in removing nanoparticles still requires in-depth research and clarification.

It is worth noting that magnetic nanoparticles have been used to treat cardiological diseases such as myocardial infarction. One of the methods of treating this type of disease is using mesenchymal stem cells (MSCs) to produce exosomes (extracellular vesicles – EVs).^{94–96} However, most cells diffuse out from the injection area immediately after cell injection into the infarcted heart, limiting their therapeutic potential. It was found that magnetic guidance of iron oxide nanoparticles MNPs–incorporated NVs following injection could improve the retention of NVs in the infarcted heart.⁹⁷ It was also demonstrated that apart from the enhanced retention, NVs derived from MNPs-incorporated MSCs contained much larger quantities of therapeutic molecules for cardiac repair (RNAs and proteins) than normal MSCs-derived NVs. Moreover, MNPs-NVs exhibited antiapoptotic, anti-inflammatory, and proangiogenic effects on cardiac fibroblasts (CFs), cardiomyocytes (CMs), endothelial cells (ECs), and macrophages, which are the primary therapeutic mechanisms of MSCs for cardiac repair in vitro.

Notwithstanding the above, studies on the biodistribution of magnetic nanoparticles in the body indicate a low level in a short time or lack of accumulation in the heart. For example, it has been reported that magnetic particle imaging (MPI) is used to track and quantify after intravenous administration of human MNPs (Ferucarbotran)-incorporated MSCs in rats.⁹⁸ The biodistribution has shown the accumulation of MNPs mainly in the liver, but no MPI signal in the heart was detected. In other studies, the authors investigated the bioavailability of carboxy dextran-coated MNPs in different organs.⁹⁹ They reported the accumulation of nanoparticles mainly in the liver, spleen, brain, and lungs and, to a small extent, also in the heart 2 hours after injection. It is worth noting that even if, shortly after administration, nanoparticles are accumulated to a small extent in the heart, their level in this organ is undetectable in the case of biodistribution measurements after a longer time, usually after 24 hours.

Brain

The blood-brain barrier (BBB) is a characteristic central nervous system (CNS) feature. The brain microvascular endothelial cells (BMVECs) line the blood vessels tightly together by tight junctions. For this reason, the penetration of various compounds is very limited. The BBB prevents the entry into the brain of various toxins and pathogens, such as



Figure 7 Structure and function of the blood-brain barrier.

bacteria, viruses, fungi and parasites, and other detrimental molecules and cells, and allows the uptake of particular nutrients and hormones only. The BBB also includes cells called pericytes and astrocytes (Figure 7A).^{100,101} Pericytes are embedded into the basement membrane of microvessels and associate closely with endothelial cells. These cells are believed to provide structural support for microvessels, influence BBB permeability, and perform immune functions by detecting and destroying harmful microorganisms. They synthesize structural and signaling proteins and contribute to the BMVECs proliferation, migration, and differentiation processes.¹⁰² The second type of cells are astrocytes that create the BBB structure contributing to the neurovascular unit. They also help recruit, among others, white blood cells to the CNS. Glial cells are immune cells that are found close to the BBB. They absorb and destroy microorganisms, another barrier against potential pathogens and toxins.

In summary, the primary function of BBB is to maintain a neutral microenvironment and protect the brain against blood-circulating microorganisms. However, BBB is semi-permeable and allows specific molecules, including, unfortunately, microbes, to penetrate through several main mechanisms.¹⁰⁰ The paracellular aqueous pathway allows water-soluble molecules to pass between endothelium cells (Figure 7B). In turn, lipid-soluble agents penetrate the lipophilic transcellular pathway. Molecules such as glucose, amino acids, and nucleosides use transport proteins, whereas insulin penetrates through receptor-mediated transcytosis. Albumin and other plasma proteins use adsorptive transcytosis. Microbes, in turn, use three main mechanisms:¹⁰³ pathogens can cross the barrier by direct endocytosis of endothelial cells, in the paracellular traversal model – through loosened tight junction, and by phagocytic penetration using transmigration within infected pathogens (the "Trojan horse" mechanism).

The BBB also prevents the delivery of therapeutics and nanoparticles to the brain for tumor diagnosis and treatment.⁴³ It turns out that if nano agents are appropriately designed, they can penetrate the BBB. Their size, charge, and coating molecules play a decisive role. For example, the functionalization of the MNPs surface with lipophilic substances improves the passive diffusion of nanoparticles into the brain parenchyma.¹⁰⁴ Also, attachment of ligands such as apolipoproteins,¹⁰⁵ transferrin antibodies and transferrin,^{106,107} α -methyl tryptophan,¹⁰⁸ and positively charged moieties have been reported to promote receptor-mediated and adsorptive-mediated transcytosis for nanoparticles to cross the BBB. It was also reported that MNPs coated with chitosan and PEG can cross the BBB.¹⁰⁹ This was related to the cationic nature of chitosan interacting with the negatively charged endothelium of the brain, as well as the small size of nanoparticles and the high lipophilicity of PEG, thus enabling penetration through the endothelial cells. Another study examined the capability of using lysophosphatidic acid (LPA) to transiently disrupt the tight junctions and allow MNPs to enter the brain.³⁸ The studies showed improved delivery efficiency of MNPs following LPA administration to the brain (approximately 4-fold vs control); moreover, no significant activation of microglia or astrocytes was observed. On the other hand, Nadeem et al¹¹⁰ reported high accumulation in the brain of the uncoated ^{99m}Tc-labelled, 10 nm-sized nanoparticles in a rabbit model. It has also been shown that dextran-coated, 30–50 nm-sized MNPs can enhance quercetin's bioavailability in the intact rat brain.³⁶

Summarizing, studies on the biodistribution of magnetic nanoparticles in mammalian organisms indicate that their accumulation in the brain is sometimes observed. The rational design of MNPs and guiding them using an external magnetic field^{111,112} allows for efficient diagnosis and drug delivery with these nanoparticles. However, the long-term biodistribution and clearance mechanisms of MNPs in the brain require further investigation.

Biodistribution of Magnetic Nanoparticles in Tumors

Cancer cells can take control of the angiogenesis process, ie, the formation of new vessels, to provide enough oxygen and nutrition for tumor growth. The formation of new blood vessels in a tumor enables its local progress and the formation of distant metastases.¹¹³ The stimulation of angiogenesis is caused by the imbalance between the factors stimulating and inhibiting this process and the most critical factor influencing the formation of new blood vessels in the vascular endothelial growth factor VEGF. In a cancerous tumor, newly formed vessels are arranged unevenly, forming numerous branches. They are characterized by changing diameters. In the walls of these vessels, there is a lack of innervation, some of the receptors, or smooth muscle endothelium (pericytes).

An essential feature of new blood vessels is their increased permeability.¹¹⁴ Leaky tumor blood vessels with pores in the endothelium with a diameter of up to several micrometers are the basis of the phenomenon known as enhanced

permeability and retention (EPR), which enables effective delivery of drugs or radioisotopes immobilized on MNPs to solid tumors.¹¹⁵ Thus, the desired nanoparticle should preferably be in the range of 20–100 nm to avoid renal clearance^{44,80} and macrophage phagocyte system (MPS)^{116,117} and be able to enter the tumors by EPR.

The use of perforated blood vessels to penetrate nanoparticles described above is a part of the so-called passive targeting mechanism. However, MNPs can be decorated by tumor-specific antibodies and directed to the specific receptor molecules found in tumor cells.¹¹⁸ There are also reports that MNPs functionalized with tumor-homing peptides can specifically bind to fibrin in tumor vessels and block blood flow, causing subsequent inhibition of tumor growth.¹¹⁹ Also guides the MNPs to the tumor sites by external magnetic field^{111,112} belongs to the so-called active targeting mechanisms.

There are many reports regarding the use of functionalized nanoparticles in the diagnostic and treatment, for example, by drug delivery to neoplastic tumors. For example, Jaiswal et al studied the efficacy of magnetic nanoparticles incorporated into poly(N-isopropylacrylamide)-chitosan nanohydrogels in inhibiting the growth of tumors in fibrosarcoma-bearing mice models.¹²⁰ The formulation was administered intratumorally to maintain the desired concentration level within the tumor mass. The study's results revealed the high efficacy of magnetic nanohydrogels in suppressing tumor growth via magnetic hyperthermia. In another study, it was reported that LTVSPWY peptide-modified PEGylated chitosan (CS)-coated magnetic nanoparticles were more actively accumulated within the area of the tumor in a short time (1 hour) as well as after 96 hours compared to PEG-CS-coated MNPs.¹²¹ The results indicated that active tumor targeting using a tumor-specific ligand was much faster and more efficient than passive targeting based on tumor permeation, uptake, and retention. Hence, LTVSPWY-PEG-CS-modified magnetic nanoparticles could be delivered to tumors by both passive and active targeting mechanisms under in vivo conditions.

On the other hand, Tietze et al used phosphate starch-coated magnetic iron oxide nanoparticles directed to the tested tumor in rabbits using an external magnetic field (magnetic drug targeting, MDT).⁶² The results showed that after intraarterial administration of nanoparticles and MDT application, MNPs accumulated in the tumor to a greater extent than after intravenous injection. In another study, MNPs were coupled to mitoxantrone.¹²² Concentrations of mitoxantrone were examined in plasma and inside the tumor 30 min after the last dose had been entirely administered. The results showed lower concentrations in the blood when a magnetic field was applied (MDT) to the tumor area, and higher doses were observed in the tumor than in the blood.

The above examples confirm that magnetic nanoparticles can be successfully accumulated in tumors using the passive targeting mechanism (increased permeability of the tumor's blood vessels) and active targeting. An active targeting mechanism by guiding MNPs using an external magnetic field increases the efficiency of nanoparticle distribution in the tumor area while reducing their concentration in the blood plasma. Also, labeling nanoparticles with peptides, antibodies, or other homing molecules significantly increases the accumulation of MNPs in specific tumors. However, it is necessary to be aware that tumors' specificity may differ depending, for example, on the site (organ) of occurrence. Therefore, the rational design of nanoparticles, the type of coating molecules, and the homing particles must be deeply studied concerning specific tumors. Moreover, nanoparticles that are expected to play a role in tumor diagnosis (Magnetic Resonance Imaging) and treatment (hypothermia, internal radiotherapy, or drug delivery) cannot be immediately captured by MPS and removed from the body. That is why it is so important to know the pharmacokinetics and biodistribution of specific nanoparticles and also, in this respect, to design nanoparticles of a specific size and coating molecules.

Factors Affecting Biodistribution of Magnetic Nanoparticles

As we recently discussed in our review, the progress in the research on the use of magnetic nanoparticles for biomedical applications has shown that their pharmacokinetics are influenced by the size, shape, charge, and, above all, surface chemistry of the nanostructures.¹²³ Similar factors determine biodistribution MNPs in body organs.

Size and Morphology of MNPs

Detailed studies confirm that the hydrodynamic diameter of nanoparticles (d_H) affects both their pharmacokinetics and biodistribution in different organs.^{33,124,125} D_H, usually measured using DLS (Dynamic Light Scattering), is the size of MNPs, including any solvent molecules attached to their surface.¹²⁶ Generally, the results of studies investigating the size

effect on the MNPs biodistribution are analogous to those found for other types of nanoparticles (for example polymeric NPs)¹²⁷ and suggest that larger MNPs are taken up faster by the liver and spleen and have shorter circulation time (and thereby half-life) in the blood. In contrast, smaller MNPs accumulate mainly in the lymph nodes and, to a lesser extent, in other organs having longer circulation time. This is confirmed by the study, which it was compared the biodistribution of Ferumoxtran-10 with the earlier reported biodistribution of Ferumoxides¹²⁸ – MRI agents with similar dextran-coating but different sizes: $d_{\rm H} = 15-50$ nm and 62–80 nm, respectively.^{129–131} In rats, 30 nm-sized Ferumoxtran-10 nanoparticles were mainly localized in the spleen: 37-46% of the injected dose (ID), liver (25% ID), and lymph nodes (5-11% ID) 24 hours after intravenous injection, whereas 80 nm-sized nanoparticles of Ferumoxides had predominant liver accumulation (83% ID at 1 h post-injection and >50% ID after 24 h).¹³² The results of these studies are in agreement with those of others where it has been studied the biodistribution of dextran-coated MNPs with sizes in the range of 33 to 99 nm.¹³³ It has been reported a similar trend: the higher distribution level of larger MNPs in the liver. In another study, the biodistribution of PEG-coated MNPs with the sizes of d_H=16.5 and 35.8 nm was analyzed.¹²⁵ The results demonstrated that both types of MNPs tended to be distributed primarily in the liver and spleen, while in other organs such as the lungs, heart, and kidneys, the presence of MNPs was found in smaller amounts. The accumulation of larger nanoparticles was slightly higher in the liver and lungs than in the case of smaller MNPs, and much less in the heart and kidneys. On the other hand, MNPs with a size of 16.5 nm achieved higher tumor uptake.

Despite the general trend outlined above, it should be considered that the pharmacokinetics, biodistribution, and elimination of nanoparticles are the result of many factors such as, among others, size, but also charge, and surface chemistry. For this reason, the results of size-dependent biodistribution may differ for a specific type of nanoparticle. Citing the detailed results as an example, the worth mentioning is the evaluation of carboxyl-coated MNPs in vivo to systematically assess their size-dependent (10, 20, 30, 40 nm MPNs) biodistribution in mice.⁴⁴ The results showed that the smallest MNPs (10 nm) showed the highest uptake in the liver, whereas the largest MNPs (40 nm) showed the highest uptake in the spleen on day 1 post-injection. The 10 nm IOMNs were also more quickly cleared from the kidney and readily penetrated the BBB. The iron content was significantly lower in the heart of mice treated with MNPs than in the control mice.

The findings regarding the influence of the MNPs' size on their biodistribution concern mainly spherically shaped nanoparticles. However, not only the size but also the shape of MNPs affect their accumulation in different organs. Studies on how the shape contributes to the biological effect of nanomaterials are limited. There are few reports on shape's impact on nanomaterials' biological performance. One of them presents magnetic mesoporous silica nanoparticles (M-MSNPs) with different aspect ratios (AR=1, 2, and 4; spheres and nanorods).¹³⁴ Biodistribution studies have shown that the shape of the nanostructures played a crucial role in the ability of M-MSNPs to accumulate in various tissues. M-MSNPs nanorods accumulated most in the spleen compared to all nanostructures tested, while the spherical M-MSNPs accumulated mainly in the liver. M-MSNPs with the highest aspect ratio in neoplastic tumors were predominantly present, while spherical M-MSNPs accumulated in the tumor the least efficiently. The authors of the discussed studies hypothesized that the significant accumulation of nanorods in the tumor was due to the larger surface area of this type of nanomaterials compared to spheres, which led to a slower clearance rate. Moreover, it is most likely related to the fact that a sizeable length-to-weight ratio translates into longer nanoparticle circulation time caused by the lesser macrophage uptake.^{42,135,136} Thus, non-spherical MNPs accumulate in the target tissues more efficiently.

More details on the biodistribution of MNPs of different sizes are provided in Table 1.

Surface Charge

The surface charge also determines biodistribution and interaction with other biological environments. Magnetic iron oxide nanoparticles with a neutral surface charge exhibit longer circulation time and reduced uptake via RES organs due to decreased opsonization. Positively charged nanoparticles are generally more easily internalized than neutral and negatively charged nanoparticles due to strong interactions with biological membranes.¹⁴² For example, Papisov et al studied the effect of surface charge on the biodistribution of two types of functionalized MNPs: the positively charged (poly-lysine)-dextran-coated MNPs and negatively charged succinate-(poly-lysine)-dextran-coated MNPs.¹⁴⁰ The positively charged MNPs showed faster clearance. They were found mainly in the liver and spleen, whereas the negative ones

Core/ Hydrodynamic Size	Coating Molecule	Animal Model	Dose	Administration Method	Technique	Main Sites of Biodistribution (Fe/g of Tissue; Fe/dl of Tissue; Fe/mm ² of Tissue)	Toxicity	Ref
10–11 /- nm	Curcumin	Mouse	5 mg/kg in 6 repeated doses	Intravenous injection	AAS ^a	Spleen and brain (≈ 0.11 mg Fe/dl); 3 days after final injection	No pathological changes in the integrity of the spleen, liver, kidney or brain	[34]
11/85 nm	Glc ^b -PMAO ^c	Mouse	0.15 mg Fe/tumor	Intratumor injection	AC magnetic susceptibility	Tumor, skin next to the tumor, spleen, liver (≈ 0.1–0.3 mg Fe/g); 30 days after iniection	-	[137
11/193 nm	Oleic acid-Pluronic	Rat	10 mg Fe/g	Intravenous injection	ICP-MS ^d	Spleen (≈ 1.2 mg Fe/g); 21 days after injection	Transient increase in oxidative stress which did not affect cellular integrity and tissue morphology	[35]
a) -/20 nm b) -/110 nm	Starch	Mouse	80 mg Fe/kg	Intravenous injection	ICP-MS	 a) Spleen (≈ 2.5 mg Fe/g); liver (≈ 1.0 mg Fe/g); b) Spleen (≈ 2.3 mg Fe/g); liver (≈ 1.6 mg Fe/g); 3 days after injection 	-	[61]
10/47 nm	Polyacrylic acid	Mouse	a) 8 mg Fe/g b) 20 mg Fe/g c) 50 mg Fe/g	Intravenous injection	AAS	 a) Spleen (≈ I mg Fe/g); liver (≈ I mg Fe/g); lungs (≈ 0.4 mg Fe/g); b) Spleen (≈ 1.7 mg Fe/g); liver (≈ 1.25 mg Fe/g); lungs (≈0.38 mg Fe/g); c) Spleen (≈ 3 mg Fe/g); liver (≈ 2.3 mg Fe/g); lungs (≈ 0.055 mg Fe/g); 24 hours after injection 	Increased hepatic lipid peroxidation at the highest dose tested (50 mg Fe/g); no significant effect in the kidneys regarding lipid peroxidation	[33]
-/100-250 nm	Phosphated starch	Rabbit	-	Intraarterial injection (close to the tumor) and magnetic targeting to the tumor	Magnetorelaxometry- SQUID ^e analysis	Liver (\approx 44% of the complete iron detected); tumor (\approx 23% of the complete iron detected)	-	[62]
5/41 nm	Uncoated, ^{99m} Tc-labelled	Mouse	100 µg Fe per mouse	Intravenous	Direct counting using an HPGe ^f detector	Liver (≈ 55% of ID ^g /g); spleen (≈ 20% of ID/g); 5 minutes after injection	-	[63]
-/40 nm	PEG ^{h, 111} In-labelled	Mouse	-	Intravenous	Assay for ⁿ C using the liquid scintillation counter	Liver (\approx 37% of the radioactivity/g); spleen (\approx 21.5% of the radioactivity/g); kidney (\approx 8% of the radioactivity/g); 24 hours after injection	-	[64]

Table I Biodistribution of Magnetic Iron Oxide Nanoparticles (MNPs) of Different Sizes and Coated with the Different Types of Molecules in Mammal Models

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12 nm/- and 15	DMSA ⁱ	Pig	a) 0.5 mg/kg	Intravenous	MRI ^j , histology, pERP ^k	Liver and spleen (majority of the injected	-	[88]
nm/-			b) 2 mg/kg			nanoparticles, dose dependent		
						accumulation); lungs [a) 12 nm-sized		
						MNPs ≈ 20 mg Fe/g, 15 nm-sized MNPs ≈		
						18 mg Fe/g; b) 12 nm-sized MNPs ≈		
						90 mg Fe/g, 15 nm-sized MNPs ≈ 50 mg/		
						g]; 5 hours after injection		
12 nm/ a) 42 nm	a) DMSA	Rat	0.44 mg Fe/kg	Intravenous	MR relaxation	a) Liver (≈ 0.5 mg Fe/g); spleen and lungs	No changes in the morphology of the	[37]
b) 49 nm	b) PEG				measurements	(≈ 0.4 mg Fe/g); b) Liver (≈ 0.42 mg Fe/g);	liver, lungs and spleen	
						lungs (≈ 0.4 mg Fe/g); spleen (≈ 0.25 mg		
						Fe/g); 3 hours after injection		
10 nm/242 nm	PAMAM ^I dendrimer	Mouse	5 mg/mL per mouse	Intraperitoneal	ICP-MS	Kidney (≈ 2.6 ppm of Fe); liver (≈ 2.0	No significant blood parameters and	[84]
						ppm of Fe); lungs and heart (≈ 1.5 ppm	renal function indicators' changes in	
						of Fe); tumor (≈ 1.2 ppm of Fe); 4	the case of chronic exposure to the	
						hours after injection	dose of 100 mg/kg; non-significant	
							increase in ALP and ALT enzymes;	
							significant increase in direct bilirubin	
							at the dose of 10 mg/kg	
-/29 nm	EDT ^m	Mouse	5 mg Fe/kg	Intravenous	ICP-OES ⁿ	Liver (≈ 80% of ID); spleen (≈ 10% of	Minimal inflammatory effects following	[38]
						ID); brain (≈ 3.6% of ID); 30 minutes	delivery of nanoparticles to the brain	
						after injection		
Nanoworms; 5	dextran	Mouse	3 mg/kg Fe	Intravenous	Magnetorelaxometry-	Lymph nodes (\approx 40% of ID/g); spleen (\approx	-	[138]
nm core, 50 nm					SQUID analysis	15% of ID/g); liver (≈ 10% of ID); 24		
length/66 nm						hours after injection		
10 nm/-	APTES°	Mouse	a) 50 mg/kg	Intraperitoneal	ICP-MS, histology	a) Spleen (≈ 0.43 mg Fe/kg); liver (≈	Transient increase in AST and LDH	[39]
			b) 100 mg/kg			0.2 mg Fe/kg); b) Spleen (≈ 0.5 mg Fe/	enzymes level; transient increase for	
						kg); liver (≈ 0.28 mg Fe/kg); 7 days	BUN; Serum TP levels within the	
						after injection	normal range; no significant damage,	
							inflammation and any pathological	
							changes in important organs	
70–80 nm/-	PEG-PAMAM	Mouse	40 mg/kg	Intravenous	CLSM ^p	Liver and spleen (≈ 0.16 mg Fe/g);	Most parameters of various cytokines	[40]
	dendrimer					lungs (≈ 0.75 mg Fe/g); heart (≈ 0.5 mg	and enzymes within the normal range;	
						Fe/g); 24 hours after injection	no significant changes in enzymes	
							reflective of inflammatory response or	
							organ toxicity	

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Table I (Continued).

Core/ Hydrodynamic Size	Coating Molecule	Animal Model	Dose	Administration Method	Technique	Main Sites of Biodistribution (Fe/g of Tissue; Fe/dl of Tissue; Fe/mm ² of Tissue)	Toxicity	Ref.
-/302 nm	PEG-modified thiolated gelatin – ¹¹¹ In-labelled	Mouse	2 μCi/200 μL per mouse	Intravenous	Radioactivity measurements	Kidneys (≈ 45% of radioactivity/g); spleen (≈ 25% radioactivity/g); liver (≈ 20% radioactivity/g); tumor (≈ 5% radioactivity/g); 24 hours after	-	[139]
10nm/-	Uncoated, ⁹⁹ ^m Tc- labelled	Rabbit	200 µg/mL per rabbit	Intravenous	SPECT	injection Heart, lung, kidney, brain (≈ 30% of concentration activity in the brain normalized over the whole-body activity concentration); after 2 hours after injection; more detailed data were not provided	-	[110]
30–50 nm/-	Dextran-quercetin	Rat	100 mg/kg	Oral	ICP-OES	Brain ($\approx 6.2 \ \mu g/g$) after 24 hours after administration; distribution to other	No histological alterations in the liver tissue	[36]
4.4–6 nm/22±6 nm	Dextran, ¹¹¹ In-labelled	Rat	2 µg/kg – 100 mg/kg	Intravenous	Radioactivity measurements	Lymph nodes: a) mesenteric, para- aortic group (\approx 30–50% of ID/g) b) popliteal, axillary groups (\approx 10–20% of ID/g) c) cervical, thoracic, other node groups (\approx 2–20% of ID/g); liver (\approx 1.5– 2% of ID/g); other tissues (< 0.2% of ID/g); 24 hours after injection	-	[140]
-/80 nm	Dextran, ⁵⁹ Fe labelling	Rat	I mg Fe/kg	Intravenous	Radioactivity measurements	Liver (\approx 82.6% of ID or \approx 6.3 of ID/g); spleen (\approx 6.2% of ID or \approx 10.9% of ID/ g); minimal amounts in other tissues; I hour after injection	-	[133]
-/110 nm	Starch	Rat	12–25 mg Fe/kg	Intravenous; magnetically targeted to brain tumors	ICP-OES and ESR ⁵	ESR results: Liver (≈ 6000 nmol Fe/g); Spleen (≈ 8000 nmol Fe/g); Lung (≈ I 500 nmol Fe/g); Kidney (≈ 6.1 nmol Fe/ g); Tumor (≈ 39.2 nmol Fe/g); Normal brain (≈ 5.6 nmol Fe/g); 50 minutes after injection	-	[141]

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	a) 10 nm/-	Uncoated	Rat	500 mg/kg	Intraperitoneal	AAS and pEPR	AAS results: a) Liver (≈ 5000 mg Fe/kg);	Histological pathological changes in	[41]
	b) 50 nm/-			3 times a week for 5			spleen (≈ 9000 Fe mg/kg); b) Liver (≈	the liver (decomplexation of hepatic	
				weeks			5000 mg Fe/kg; spleen (≈11,500 mg Fe/	tubules, cytolysis observed in some	
							kg)	places); extensive deposits of lumps of	
							pEPR results: a) Liver (≈ 2500 mg Fe/kg);	iron-containing pigment in the red	
							spleen (≈ 7000 mg Fe/kg); b) Liver (≈	pulp of the spleen	
							3200 mg Fe/kg); spleen (≈ 8000 mg Fe/kg)		
							after 5 weeks; distribution to other		
							organs has not been studied		

Note: Dashes in some of the table cells indicate that the data was not available in the source articles.

Abbreviations: ^aAAS, Atomic Absorption Spectroscopy; ^bGlc, glucose; ^cPMAO, poly(maleic anhydride-alt-1-octadecene); ^dICP-MS, Inductively Coupled Plasma, Mass Spectroscopy; ^eSQUID - Superconducting Quantum Interference Device; ^fHPGe, High Purity Germanium; ^gPEG, polyethylene glycol; ^hID, injected dose; ⁱDMSA, dimercaptosuccinic acid; ⁱMRI, Magnetic Resonance Imaging; ^kpERP, particle Electron Paramagnetic Resonance; ⁱPAMAM, poly(amidoamine); ^mEDT - ethylenediamine triacetate; ⁿICP-OES - Inductively Coupled Plasma Optical Emission Spectrometry; ^oAPTES, (3-aminopropyl)triethoxysilane; ^pCLSM, Confocal Laser Scanning Microscope; ^qMPI- Magnetic Particle Imaging; ^rSPECT, Single-Photon Emission Computed Tomography; ^sESR, Electron Spin Resonance.

accumulated in lymph nodes, thus showing more similar biodistribution to the nearly neutrally charged dextran-coated MNPs. Another study found that both negatively and positively charged MNPs showed increased liver uptake compared to neutrally charged MNPs.¹³³ On the other hand, Sun et al studied the biodistribution of MNPs coated with negatively charged ethylenediaminetetraacetate (EDT) as the most likely candidates for the BBB crossing to enhance the delivery of MNPs in the brain.³⁸ The selection of negatively charged MNPs-EDT was based on several previous findings. Firstly, negatively charged nanoparticles have improved colloidal stability¹⁴³ and lower levels of non-specific cell association compared to positively charged MNPs.¹⁴⁴ Secondly, MNPs-EDT in vitro cell culture models were significantly more permeable than positively charged MNPs following osmotic disruption and magnetic field applications.¹⁴⁵

Coating Molecules and Protein Binding

Coating molecules on the MNPs surface affect their size and charge, influencing their biodistribution described in previous chapters. However, it is essential to consider the complexity of the process as surface chemistry also affects the binding protein profile, and the binding proteins influence biodistribution. A binding protein can cause additional changes in nanoparticle size and charge, affecting the internalization process of MNPs into macrophages and the overall distribution in the body.^{146–148}

Specific bound proteins, such as IgG, complement factors, and fibrinogen, allow macrophages of RES to recognize nanoparticles and promote phagocytosis more easily.^{149,150} The particles coated with these proteins tend to accumulate in the liver and spleen. For imaging and drug delivery to body areas other than RES organs, MNPs must have a longer circulation time in the blood and reduced flow through the RES organs. The polymers that give the MNPs an anti-fouling character and enable their increased uptake to the site of interest are different variations of polyethylene glycol (PEG) or poly(ethylene oxide) (PEO)-containing chemical groups.^{151–154} Generally, PEGylation is the most common method of masking nanoparticles from immune recognition.¹⁵⁵ It has been shown to decrease interactions of nanoparticles with blood proteins and help avoid recognition by RES, thus prolonging circulation time in blood and making it easier to reach organs other than the liver and spleen. For example, Liu et al¹⁵⁶ developed single-core Fe₃O₄ MPI tracers coated with covalently bonded PEG-silane brushes. The newly tailored tracer had MPI sensitivity about three times higher than the commercial tracer Ferucarbotran and 3 minutes after injection in the mice model, the tracer was seen to be distributed throughout the body, with the most significant signal corresponding to the heart. There are also studies showing that PEG-coated can effectively distribute to the tumors. For example, MNPs with a crosslinked starch and PEG coating showed increased circulation time (up to 12 h) and distribution in the tumor.¹⁵²

PEG and many other coating polymers and molecules and their effect on biodistribution in the body were tested. For example, a high signal from Pluronic-oleic acid-MNPs was reported at the tumor site on the second day after administration.¹⁶ Jain et al³⁵ developed iron oxide MNPs coated with both oleic acid (OA) and Pluronic that can be loaded with high doses of hydrophobic drugs¹⁵⁷ and studied their biodistribution in rats. The maximum iron level was observed in the liver after 6 h (55% of the total injected iron), which was only 20% after one day. These MNPs have been accumulated in other organs, mainly in the spleen, lungs, kidneys, heart, and brain. Another study determined the concentration of MNPs coated with oxidized starch and their degradation products in Kupffer, endothelial and parenchymal cells of the rat liver.¹⁵⁸ Even though the nanoparticles had a small size and quite prolonged circulation time, approximately 80% of the injected MNPs were taken up by the liver within 24 h after single-dose administration. It is worth mentioning that nanoparticles exhibited negative zeta potential because of the coating by the oxidized starch. Therefore it can be assumed that the uptake of the tested nanoparticles was mediated by scavenger receptors expressed in Kupffer cells and liver endothelial cells, binding ligands with clusters of negative charge. Ikeda et al¹⁵⁹ synthesized activatable fluorescence agents of dextran-coated MNPs at various molecular ratios and hydrodynamic diameters (34.1-35.9 nm). The most significant signal intensity in biodistribution studies was observed in the liver, followed by the spleen. When an excessive amount of nanoparticles were administered, a portion of the circulating MNPs was taken up by macrophages in the liver and spleen, and the excess was detected in other tissues, such as the lungs and adipose tissue, where macrophages are substantial.⁵⁰ On the other hand, there were developed theranostic hybrid MNPs with carbon shells and a hydrodynamic diameter of about 16 nm.¹⁶⁰ Biodistribution studies have shown that nanostructures were mainly accumulated in the liver through the RES system. However, the carbon-shell MNPs were also detected in the

kidneys, spleen, lungs, heart, and tumors. There were also developed dextran-coated ultrasmall iron oxide nanoparticles coated with peptides containing L-arginine-glycine-aspartate (Arg- Gly-Asp, RGD)¹⁶¹ and aspartic acid-glycinearginine (Asn-Gly-Arg, NGR)¹⁶² as potential double-targeting agents in tumor angiogenesis magnetic resonance imaging (MRI).¹⁶³ The peptides bind specifically on the endothelial cells' surface in tumor neovascularization resulting in the regulation of angiogenesis, tumor growth, and metastasis.^{164–166} The number of nanostructures in tumor tissues increased 1.83 times compared with the control group. However, they also accumulated in the kidney, liver, and spleen. Salimi et al⁸⁴ synthesized MNPs coated with the fourth generation (G4) of polyamidoamine (PAMAM) dendrimer and the amino groups of dendrimer molecules were next conjugated with mPEG.^{167,168} At 24 h after injection in the mice model. the maximum amount of the nanostructures was found in the lung and kidney tissues. The results showed decreased PAMAM-PEG coated nanoparticles' biodistribution in all tissues 8 h after administration. The iron level was reduced by about half from the initial dose except for the lungs, where the iron content decreased by 14%. The authors explained that this phenomenon could be the return of nanoparticles to the bloodstream and then uptake to the spleen or lymph nodes. At the same time, the lung immune system acted as the main barrier against PAMAM-PEG-coated MNPs and prevented them from returning to the bloodstream and other organs.¹⁶⁹ It also evaluated the biodistribution of three types of MNPs that were coated with a carboxyl group, carboxyl group followed by PEG coating, and carboxyl group followed by BSA coating, all in the range of sizes 54–62 nm.¹⁷⁰ The studies conducted in rats have shown that the iron content in the kidneys, intestine, and testis did not show a significant difference compared with the control group. PEG-MNPs and COOH-MNPNs accumulated in the liver and lung. BSA-coated MNPs showed more accumulation in blood than PEG-MNPs and COOH–MNPs, which indicated that naturally occurring compounds, such as BSA, were more suitable for surface modification. After 24-day recovery, BSA-MNPs had longer-term retention time in blood, and PEG-MNPs seemed to have more accumulation ability in various organs than MNPs coated with BSA. Thus, the authors demonstrated that surface modifications played an essential role in the biodistribution of magnetic nanoparticles.

More details on the biodistribution of MNPs with different coatings are provided in Table 1.

The Effect of MNPs Vectorization

Detection, drug delivery, and imaging (MRI) by MNPs are typically limited to regions rich in the mononuclear phagocyte system, such as the liver and spleen. However, using an external magnetic field allows directing of magnetic nanoparticles to specific body sites. Another approach to achieve targeted drug delivery, especially to tumors, consists of grafting functional groups on the surface of MNPs with ligands that recognize specific receptors or biomarkers. Typically, these ligands are oligosaccharides,¹⁷¹ peptides and proteins^{172–179} (eg, insulin, RGD peptides, transferrins), antibodies or their fragments,^{180–183} mimetics,¹⁸⁴ as well as vitamins such as folates that interact with tumor cell receptors.^{185,186} This facilitates the biodistribution of MNPs into specific body regions. Furthermore, the antibodies labeled with low doses of radionuclides and immobilized on MNPs are used in radioimmunotherapy (RI).¹⁸⁷ Additionally, fluorophores anchored on the surface of nanoparticles are useful in imaging and diagnosis by using various diagnostic techniques.

More specific information concerning selected targeting agents immobilized on functionalized MNPs is presented in Table 2.

Technical Aspects

Some technical aspects, such as the way of administration, can influence the biodistribution of MNPs in the body. The most common is an intravenous injection. However, several reports describe using MNPs coated with organic shells in oral administration. One of them concerns using MNPs as contrast agents for the gastrointestinal tract.¹⁸⁹ In another study, a novel drug delivery system was reported composed of layer-by-layer milk protein casein-coated iron oxide nano-particles with high stability in the stomach and enzyme-response release in the small intestine.¹⁹⁰ It was also reported using a nanocomposite, phytic acid-chitosan-magnetic iron oxide nanoparticles in treating colon cancer.¹⁹¹ However, the studies on the biodistribution of MNPs after oral administration are very limited. Generally, MNPs must pass the transport barrier of the intestinal epithelium, and the following can cross the liver sinusoids and then gain entry to the central blood circulation system. Thus the liver is the major accumulation and clearance organ in the oral administration of MNPs, whereas the nanoparticles remaining in the digestive system are extracted through the feces.¹⁹²

Type of	Specific Targeting	Target	Coating	Application	Ref.
Agent	Agent				
Proteins	Transferrin	Transferrin	DMSAª	Brain glioma imaging	[175]
		receptor	Chitosan/Doxorubicin		[1/6]
	Lastafamin (family of		Dextran-spermine	Drug delivery by BBB	[1//]
	transferrins)	receptor-related	FILAO	Drug deliver by BBB (gloma cells)	[170]
		protein			
Peptides	GRGD ^a -peptide	ανβ 3 integrin	3-Aminopropylsilane	Not specified	[172]
	CRGD ^a -peptide		PEG®	Imaging and drug delivery	[173]
	Insulin	Human	Sodium oleate	Drug delivery	[179]
		fibroblasts			
		receptors			
Antibodies	SM5-1 monoclonal	Hepatocellular	Dextran	Targeted contrast agent for diagnosis	[180]
	antibody	carcinoma cells		of hepatocellular carcinoma cells	
	Anti-transferrin	Transferrin	MNPs enveloped in	Drug delivery to the brain	[181,182]
	receptor antibody	receptor	a PEGylated bilayer to form		
	OX26		liposomes		
	Anti-EGF ^f	EGF-receptor	PEG	Detection of lung cancer using MRI	[183]
	monoclonal antibody				
Other	Sialyl Lewis ^x (sLe ^x)	E-selectin	Dextran	MRI diagnosis of inflammation	[184]
molecules		(leukocytes)			
	Folic acid	Folate receptor	APTES ^g /PEG	Chemotherapy and hyperthermia	[185]
			Polyethyleneimine	Tumor targeting MRI	[186]
	Hyaluronic acid	CD44 antigen	Polymeric micelles using	MRI and photothermal-	[188]
			hyaluronic acid -C ₁₆	chemotherapy	
			copolymers		

 Table 2 MNPs Decorated with Targeting Agents

Abbreviations: ^aDMSA, dimercaptosuccinic acid; ^bBBB, Blood Brain Barrier; ^cPMAO, Poly(maleic anhydride-alt-1-octadecene); ^dRGD-peptides family, L-Arg-Gly-L-Asp; ^ePEG, Polyethylene glycol; ^fEGF, Epidermal Growth Factor; ^gAPTES, (3-Aminopropyl)triethoxysilane.

In the case of subcutaneous or intratumoral administration, as a general rule, the injected nanoparticles need to be small (60 nm or less).¹⁹³ These small particles injected locally infiltrate the interstitial spaces around the injection and are absorbed by the lymphatic capillary system.¹⁹⁴

Regarding the intraperitoneal administration, it was tested biodistribution in a rat model of naked Fe_3O_4 nanoparticles of three sizes: 20, 50, and 1000 nm using, among others, Atomic Absorption Microscopy (AAS).⁴¹ The results showed a high accumulation of 10– and 50 nm-sized MNPs in the spleen, followed by the liver. In comparison, micrometric particles (1000 nm) were distributed only slightly and statistically insignificantly compared to the organs of control rats.

Nanoparticles administered by inhalation or intratracheally are introduced into the alveolar spaces and mainly used for diagnosing and treating lung diseases.^{169,195} For example, studies in mice have shown that inhaled MNPs accumulate in the central region of the lung about 2.5 times more efficiently than in the peripheral zones of this organ. MNPs are phagocytized by macrophages in the alveolar spaces, and their by-products are released into the pulmonary lymphatics.^{196,197}

Another technical factor with a significant influence on the biodistribution of nanoparticles is the use of an external magnetic field to guide MNPs to specific sites, eg, tumors. For example, it investigated the biodistribution of¹²² Iod-labelled ferrofluids (about 100 nm MNPs coated with hydrophilic starch) in vivo in rabbits.¹⁹⁸ The results of studies showed that ferrofluid concentration was dependent on the strength of the magnetic field. In addition, after using the focused external magnetic field, the scintigraphically detected¹²² Iodine signal after intra-arterial administration to the tumor in the left femoral artery was significantly higher than administration without the magnetic field. After 10 and 60

minutes after administration of the nanoparticles, 77% and 22% of their initial activity in the tumor area were found, respectively, in the case of the application of an external magnetic field, while only 40% and 11%, respectively, in the absence of a magnet. In another study, the histology method investigated the effects of an external magnetic field on the biodistribution of MNPs in a rat model.¹⁹⁹ The dextran-coated Fe_3O_4 MNPs (250 nm) functionalized with the -COOH group were administered intravenously. Biodistribution studies showed accumulation of MNPs mainly in the liver and spleen in the case of the control (without magnet). When the magnetic field was applied to the right kidney, MNPs were retained over both kidneys with more of the right side. Other organ tissues close to the right kidney (ie, stomach, small intestine, and colon) did not have significant retention of MNPs.

There was also studied on the effect of anesthesia on MNPs' biodistribution after MNPs' intravenous injection.²⁰⁰ For example, animals (murine) were anesthetized with isoflurane (0.5% in oxygen) or by intraperitoneal injection with a mixture of ketamine and xylazine. Then, dimercaptosuccinic acid-coated MNPs were intravenously injected. The number of MNPs in the liver and lung tissue was measured using the Alternating Current Magnetic Susceptibility method. The results have shown that the accumulation of nanoparticles in the liver was not affected by the method of anesthesia. However, the number of particles that reached the lungs for inhaled isoflurane was three times less than for the intraperitoneally injected anesthetic. In conclusion, this parameter (the type of anesthesia) must be considered when comparing different studies on MNPs distribution.

The most important factors affecting the biodistribution of magnetic nanoparticles are summarized in Figure 8.

Biodegradation and the Fate of MNPs in the Organisms

Iron oxide magnetic nanoparticles can be removed from the body by renal clearance mechanism²⁰¹ or by MPS (phagocytic cells in the blood, tissues, lymph nodes, etc.).¹²⁵ Generally, the healthy human body contains 65% iron as hemoglobin, 4% as myoglobin, and 15–30% and 0.1% as ferritin and transferrin, respectively.²⁰² Ferritin is a protein that complexes Fe^{3+} iron ions and stores them in the liver, keeping the iron in an accessible and harmless form. In contrast, transferrin is a glycoprotein that regulates the concentration of iron ions in the blood plasma and transports them to the tissues.²⁰³ The mechanism of the biodegradation of magnetic nanoparticles is believed to be analogous to the metabolism of ferritin, which is digested by lysosomal enzymes to release iron ions.³² The excess of iron (ferrous Fe^{2+} or ferric Fe^{3+} products) is then regulated by various clearance mechanisms of the body, namely, it is integrated into either the storage or utilization pathway.^{125,204}



Figure 8 The main factors affecting the biodistribution of MNPs.

Intravenously administered magnetic nanoparticles are captured by macrophages, mainly of the liver and spleen, and then digested in the acidic environment of the lysosomes.^{205,206} Degradation of MNPs occurs faster in the liver macrophages than in the spleen, as the former contains more iron-storing proteins.²⁰⁷ The degradation rate depends on the material covering the nanoparticles²⁰⁸ (hydrophilic substances favor faster degradation) as well as the dose of nanoparticles.²⁰⁹ The released iron binds with apoferritin present in the cytoplasm of macrophages to form a ferritin.⁷⁹ After the iron is transported from ferritin to apotransferrin, transferrin is formed.²¹⁰ Transferrin circulates through the body supplying iron to various tissues, such as muscles, where myoglobin is biosynthesized, or bone marrow, where iron forms hemoglobin, which gets stored in the red blood cells (RBC).⁷⁹ Iron ions can be incorporated in the RBC hemoglobin as ferrous (Fe²⁺) or ferric (Fe³⁺) ions, albeit only ferrous ions are active in oxygen transport. Nevertheless, methemoglobin reductase can reduce iron ions in the third oxidation state to Fe²⁺ ions.²¹¹ Therefore, there may be several alternative metabolic pathways for different forms of iron (i. a. FeO, Fe₂O₃, Fe₃O₄), which requires in-depth research.

The senescent RBCs are taken up by the red pulp of the spleen, then phagocytosed by macrophages and converted to ferritin and subsequently transferrin, which redirects iron to the bone marrow to form new erythrocytes, or to the liver, where iron is stored in hepatocytes as ferritin.¹⁵⁸ A certain fraction of hemoglobin is also converted to bilirubin, then excreted in urine and bile. If the body's iron content exceeds the available apoferritin, the iron forms small insoluble aggregates called hemosiderin in the liver cells, slowly releasing iron ions.²¹²

MNPs can also be excreted via urine; however, nanoparticles must be small enough to utilize renal clearance, with a diameter below 10–15 nm.²⁰¹ For example, MNPs with a size in the range of 1–5.5 nm are assumed to be renalclearable, as they can readily pass through the glomerular filtration membrane.

Nanoparticles designed for MRI or drug delivery to specific regions of the body, such as a brain tumor, must be able to cross the blood-brain barrier.²¹³ After fulfilling their function, the MNPs should be quickly cleared from the brain as the accumulation of nanoparticles could lead to neurotoxicity. Gu et al²¹⁴ investigated the clearance of MNPs coated with reconstituted high-density lipoprotein (HDL) and with poly(ethylene)-glycol-b-poly(lactic acid) copolymer (PEG-PLA) from the brain. They reported that the studied nanostructures were easily and rapidly cleared through the perivascular glymphatic pathway. This pathway involves astrocytes - The giant glial cells, which generally perform many functions, including participation in nerve transmission, ammonia and creatine transformations, regeneration, phagocytosis processes, and removing soluble proteins and metabolites from the brain.

Current Techniques for Analyzing the Biodistribution of Nanoparticles

The biodistribution of nanoparticles in the body means the place of their accumulation in various tissues and organs, the way they interact, their therapeutic effect, their residence time, and elimination from tissues. Knowing the biodistribution after in vivo administration of nanoparticles is essential primarily because of the target organs to which MNPs are to be delivered, but also because of undesirable accumulation and toxicity in other tissues. Therefore, the nanoparticles used in the clinic must first be thoroughly tested for biodistribution in mammalian models. Several different techniques are used for this purpose.

One of the most popular techniques for evaluating nanoparticle biodistribution is histology,²¹⁵ also known as microscopic anatomy or microanatomy. The technique involves the histological examination of tissues to determine the association of nanoparticles with the cellular microenvironment under a microscope, usually light and fluorescent. Selected tissue fragments are harvested at set time points and traditionally stained with Prussian blue dye. For example, Pham et al²¹⁶ studied the biodistribution of ultrasmall MNPs injected intraperitoneally into mice by measuring iron-positive areas in tissue sections of organs stained with this dye. In terms of advantages, histology is an inexpensive technique to study the biodistribution and cannot distinguish between individual nanoparticles in a tissue. Moreover, the technique is sensitive to human error in sample preparation and analysis techniques. Regarding fluorescence imaging, labeling MNPs with fluorescent dyes may influence their physicochemical properties and, thereby, their in vivo behavior.²¹⁷

Another standard laboratory method is Liquid Scintillation Counting (LSC).²¹⁵ The technique quantifies the radioactivity of low-energy radioisotopes, mostly β -emitting and α -emitting isotopes.²¹⁸ LSC sample requires a mixture of the aromatic organic solvent that absorbs radioisotopic energy and scintillators ("fluors") to produce detectable light pulses. The energy released from radioactive particles excites the aromatic solvent molecules, producing excited states of electrons by scintillator molecules. Their decay to the ground state causes producing a light pulse that is characteristic of the scintillator. The emitted light is detected by the liquid scintillation counter's photomultiplier tube (PMT). For in vivo biodistribution studies, nanoparticles are labeled with isotopic markers. Organs are then harvested and prepared for LSC. The technique is sensitive, specific, and quantitative. However, an essential issue in this method is the separation of the excess of the free isotopes that did not incorporate into the nanoparticles. Moreover, LSC determines only biodistribution at the tissue or organ level but does provide any information on the specific cellular interactions.

Electron microscopy analysis provides detailed information on the biodistribution of nanoparticles under very high magnification.²¹⁹ There are different types of electron microscopy techniques, such as Transmission Electron Microscopy (TEM), Scanning Electron Microscopy (SEM), and variation of these two techniques, for example, Scanning Transmission Electron Microscopy (STEM).²¹⁵ In these techniques, the electron beam is focused by electromagnetic lenses and directed at the sample. In the case of TEM, the beam passes through the sample, while in the SEM technique, the electron beam scans the sample's surface, and the signal intensity is mapped synchronously with the scan, which allows for an enlarged image.²²⁰ Electron microscopy provides detailed information on the biodistribution of nanoparticles, but this information is limited to a certain number of ultra-thin tissue sections. On their basis, the approximate biodistribution of MNPs in the whole organ is estimated.²¹⁵ It is worth emphasizing, however, that these techniques enable the visualization of the accumulation of nanoparticles in individual cells and organelles. The disadvantage of these techniques is their cost and time consumption. Moreover, they are used more often to determine the cellular association of nanoparticles in vitro. In contrast, the studies using electron microscopy to evaluate nanoparticle biodistribution after in vivo administration are limited.^{219,221–223}

The Laser Scanning Confocal Microscope (LSCM) is used in biomedical research. Confocal microscopy is a variant of light microscopy that allows obtaining images of high magnification and resolution. The LSCM uses a laser as the light source, sensitive tube detector photomultipliers (PMTs), and a computer to control the scanning mirrors and to collect and display the images.²²⁴ The diffraction-limited point in the sample is focused by the objective lens and scanned across by the scanning device, and thereon the point of light is detected by the PMT. The output from the PMT is embedded into the image by the computer. The LSCM method was used, among others, to determine the biodistribution of PAMAM dendrimer conjugated MNPs in mice⁴⁰ and folic acid conjugated MNPs for drug delivery (mice).²²⁵

Magnetic Resonance Imaging (MRI) uses strong magnets to generate a magnetic field. Under its influence, the protons in the body align with the field. MRI measures the spin magnetization of polarized protons and their respective longitudinal (T_1) and transverse (T_2) relaxation rates, resulting in high-resolution images. Contrast agents increase the speed at which protons realign and shorten the relaxation rates. Paramagnetic contrast agents reduce the longitudinal (T_1) relaxation time, which results in brighter images, whereas superparamagnetic contrast agents (MNPs) shorten the transverse relaxation rate (T_2), resulting in darker images.^{226,227} On the other hand, it has been reported that MNPs smaller than 5 nm have decreased magnetic moments, which causes a strong T_2 suppression effect. Consequently, ultrasmall MNPs can be exploited as T_1 agents.²²⁸ Generally, MRI is a direct and non-invasive technique that can determine nanoparticle biodistribution at the tissue and organ level and has unlimited penetration.²¹⁵ However, it is a relatively costly technique. Additionally, this technique can need better sensitivity.

Also, Magnetic Particle Imaging (MPI) is a diagnostic in vivo imaging method designed to image the amount and location of superparamagnetic nanoparticles in biological tissue. It makes use of the nonlinear magnetic response of the MNPs.²²⁹ The MPI technique allows direct detection of nanoparticles, which results in greater sensitivity than magnetic resonance imaging (MRI).²³⁰ Moreover, the intensity of the image obtained by MPI is directly related to the concentration of detected MNPs, so the technique enables the quantitative study of nanoparticles. In addition, MPI is not as expensive as MRI because it does not require an expensive superconducting magnet.²³⁰ MPI exploits the combination of an alternating magnetic field gradient and a static magnetic field gradient for real-time. During MPI scanning, the nanoparticles are magnetized by a radio frequency (RF) magnetic field, and their time-varying magnetization induces

a time-dependent voltage across the inductively coupled receiver coil.^{231,232} To sum up, MPI offers quantitative, high-contrast, and high-resolution images.

Another technique used for determining the biodistribution of MNPs is Single Photon Emission Computed Tomography (SPECT).²³³ It is a non-invasive technique that produces three-dimensional images of the body by detecting gamma rays emitted from radioactive substances (which the nanoparticles must be coated with) that become localized and are taken up by specific tissues.^{234,235} As the isotope decays, it emits gamma rays picked up by detectors (direct measurement). The most common radioisotopes used for SPECT imaging include ^{99m}Tc, ¹¹¹In, and radioiodine.^{110,234,236} The biodistribution of nanoparticles can be assessed in real-time, and tissue penetration limits do not restrict the technique.²¹⁵ Conversely, it is a relatively costly technique requiring ionizing radiation exposure.

Magnetic Relaxometry (Magnetorelaxometry, MRX) can detect and locate targeted magnetic nanoparticles noninvasively without ionizing radiation. In the first phase of the magnetorelaxometric measurement, an external magnetic field of a defined strength and duration partially aligns the MNPs' moments in the direction of the field. In the second phase, just after the external field is switched off, the fading magnetic field from the decay of the sample is detected by sensitive sensors such as a superconducting quantum interference device (SQUID).²³⁷ By this method, very sensitive detection of MNP amounts down to a few nanograms is feasible in vitro or ex vivo.^{237,238} For MRX to be used in biological and medical applications, the particles must possess definitive size ranges (on the order of tens of nanometers) and have high magnetic moments.²³⁹ This method separates nanoparticles bound to objects (for example, cells) from those that rotate in a freeway in a suspension liquid.²⁴⁰ The technique's sensitivity depends on the net magnetic moments induced in the single-domain MNPs by the magnetizing pulse, the number of MNPs bound in the sample, and the sensitivity of the magnetic field sensors (such as SQUID).²³⁹

Electron Paramagnetic Resonance (EPR) spectroscopy, called also Electron Spin Resonance (ESR) spectroscopy can be compared to NMR (nuclear magnetic resonance) spectroscopy as both techniques use the interaction of magnetic dipoles with an applied magnetic field and electromagnetic radiation of the appropriate wavelength.²⁴¹ However, NMR relies on splitting nuclear spin states in a magnetic field, whereas EPR/ESR on splitting electronic spin states. Electron magnetic dipole arises from one or more unpaired electrons. Therefore, not all molecules are "EPR/ESR active", as the most stable molecules have a closed electronic shell. Thus, EPR/ESR spectroscopy is used to study chemical species with unpaired electrons, including paramagnetic (such as Fe(III) – transferrin), ferrimagnetic and ferromagnetic (eg, magnetite), and antiferromagnetic (eg, ferritin) materials.^{242,243} Both endogenous iron-protein complexes (such as hemoglobin, transferrin, ferritin, and hemosiderin) and the exogenous MNPs can be detected and distinguished with EPR/ESR. Although the technique is appropriate to study iron oxide nanoparticles and free radicals in the biological samples,^{244,245} it has not been widely utilized for biodistribution studies.^{242,243} This limited use could be attributed to the technical challenge of loading tissues into a long (190 mm), narrow (3 mm) EPR/ESR tube for sample analysis.¹⁴¹

Another technique often used for determining the biodistribution and concentration of iron contained in MNPs is Atomic Absorption Spectroscopy (AAS).^{33,34,41,216} AAS is an analytical technique used to determine how many specific elements are in a sample. It uses the principle that atoms and ions can absorb light of a specific wavelength. The atom absorbs this particular wavelength of light. The electrons in the atom go from the ground state to the excited state, and the amount of absorbed light is measured. In this way, the concentration of an element in the sample can be calculated.^{246,247} AAS is a well-known technique. It is easy to use and provides high accuracy and sensitivity.^{248,249} Importantly, AA spectrometry does not cost much to operate. The AA spectrometer can measure samples from low ppm to percentage levels over a wide concentration range. However, it can be challenging to determine elements present in very high concentrations.^{248,249} Moreover, AAS does not allow for the differentiation of endogenous and administered iron.

The Alternating Current Susceptibility (ACS) is also used in the biodistribution determination of MNPs.¹³⁷ In this technique, the sample is subjected to an oscillating external magnetic field, and during this process, the magnetic moment of the sample is measured.²⁵⁰ A typical measurement system consists of a permanent magnet providing a static field and a modulating coil that generates an alternating current magnetic field and produces a time-dependent magnetic moment in the sample. Only the signal induced by the magnetized sample is measured, and the background is subtracted. The ACS technique distinguishes the applied nanoparticles in the imaged tissue fragment from endogenous iron.²⁵¹ Additionally, maximum contrast for specific types of nanoparticles can be obtained by selecting specific frequencies. Moreover, ACS

provides information on the surface chemistry of MNPs and their agglomeration.²⁵² It has also been shown that the ACS technique can monitor the degradation of MNPs in biological media.²⁵³

Inductively Coupled Plasma (ICP) techniques coupled with Atomic Emission Spectroscopy (ICP-AES)^{254,255} or Mass Spectroscopy (ICP-MS)^{256,257} also are applied for determining of MNPs distribution. These techniques use electromagnetic induction at high temperatures, generating argon plasma, thus breaking molecular and ionic bonds in the sample. With ICP-MS, the sample is ionized, and the mass spectrometer records the mass-to-charge ratio. In the case of ICP-AS, the sample atoms are excited by plasma, and electromagnetic radiation is quantified by a spectrometer. Even though these techniques have been reported to detect iron at low concentrations, the main disadvantages of these methods are their destructive nature and the fact that they do not differentiate between endogenous and administered iron.¹⁴¹

Conclusions

The review describes the biodistribution in the body of magnetic iron oxide nanoparticles used in medicine. Thanks to their biocompatible structures MNPs are increasingly translated from laboratory research to medical applications such as drug delivery, internal radiotherapy, MRI, or hyperthermia.

In addition to their primary supermagnetic properties, MNPs also have a large specific surface area, which allows the immobilization of therapeutic agents such as drugs or radioisotopes. By utilizing functionalized and specifically decorated magnetic nanoparticles, it becomes feasible to achieve dual targeting. These nanoparticles can be directed toward specific target places in the body supported by an external magnetic field as a guiding mechanism. Furthermore, the focused application of a magnetic field allows for the limitation of MNP toxicity, thereby reducing the total amount of nanoparticles within the body.

The biodistribution of nanoparticles in the body is a parameter that determines the place and degree of the administered nanoparticle accumulation in the body. The main factors influencing the biodistribution of MNPs are their size, shape, surface charge, covering molecules and specific vectors, interaction with proteins, the route of administration, and the use of an external magnetic field. In general, biodistribution is the product of all these factors. Uncontrolled proteins attachment to MNPs (opsonization) can lead to various adverse effects, including rapid clearance from the bloodstream, hindered targeting capability, reduced bioavailability, and potential toxicity induction.

Due to the mononuclear phagocyte system, detection, drug delivery, and imaging of MNPs are typically limited to the liver, spleen, and lymph nodes. To maximize the enhanced permeability and retention (EPR) effect, it is crucial to consider the overall sizes of injected surface-functionalized iron oxide nanoparticles. Multiple studies have confirmed that nanoparticles with a size of approximately 50 nm exhibit optimal cellular uptake and slower opsonization. Larger nanoparticles exceeding 100 nm are eliminated by the reticuloendothelial system (liver and spleen), whereas nanoparticles smaller than 10 nm are cleared from the bloodstream through renal clearance.

However, applying an external magnetic field enables the targeted delivery of MNPs to specific body sites. There are many reports that the proper design of nanoparticles allows these structures to accumulate in other organs, such as the kidneys, lungs, heart, and brain. A particular challenge is to design MNPs (surface chemistry, size) to cross the BBB barrier in the brain.

Moreover, attaching ligands recognizing specific receptors or biomarkers to the functional groups on the biocompatible and biodegradable surface of MNPs facilitates the biodistribution of MNPs in specific body regions, particularly to tumor areas. The MNPs are surface coated with dextran or its derivatives for clinical application. Hydrophilic polymers, such as FDA-approved polyethylene glycol, polyphosphoesthers, and zwitterionic polymers, could also be considered.

The ligands that recognize receptors on the cells include oligosaccharides, peptides, proteins (such as insulin, RGD peptides, and transferrins), antibodies or antibody fragments, mimetics, and vitamins like folates that interact with tumor cell receptors. It has been shown that the degree of accumulation in tumors is significantly increased by applying an external magnetic field additionally.

Knowing biodistribution lets us determine if the designed nanosystems target the desired tissues and organs for therapeutic purposes. On the other hand, biodistribution studies indicate undesirable accumulation that can lead to the potential toxicity of MNPs. Therefore, studies on the biodistribution of MNPs are particularly important for clinical applications. Addressing these limitations is crucial to further enhance the efficacy and safety of MNPs as drug delivery carriers.

Author Contributions

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis, and interpretation, or all these areas; took part in drafting, revising, or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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