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Magnetic particle imaging: tracer development and the biomedical applications of a radiation-free, sensitive, and quantitative imaging modality

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Magnetic particle imaging (MPI) is an emerging tracer-based modality that enables real-time three-dimensional imaging of the non-linear magnetisation produced by superparamagnetic iron oxide nanoparticles (SPIONs), in the presence of an external oscillating magnetic field. As a technique, it produces highly sensitive radiation-free tomographic images with absolute quantification. Coupled with a high contrast, as well as zero signal attenuation at-depth, there are essentially no limitations to where can be imaged within the body. These characteristics enable various biomedical applications of clinical interest. In the opening sections of this review, the principles of image generation are introduced, along with a detailed comparison of the fundamental properties of this technique with other common imaging modalities. The main feature is a presentation on the up-to-date literature for the development of SPIONs tailored for improved imaging performance, and developments in the current and promising biomedical applications of this emerging technique, with a specific focus on theranostics, cell tracking and perfusion imaging. Finally, we will discuss recent progress in the clinical translation of MPI. As signal detection in MPI is almost entirely dependent on the properties of the SPION employed, this work emphasises the importance of tailoring the synthetic process to produce SPIONs demonstrating specific properties and how this impacts imaging in particular applications and MPI’s overall performance.

1. Introduction

MPI is a recently developed tracer-based modality which has emerged as a promising diagnostic and therapeutic tool with wide ranging potential applications. It can generate 2D projection images or true 3D tomographic images that are easily interpretable, stemming from their ‘positive contrast’, a trademark of other tracer-based techniques like positron emission tomography (PET), single-photon emission computed tomography (SPECT), and optical imaging techniques. The tracer employed by MPI are SPIONs. MPI utilises a gradient field with strong gradients and weak field strengths, and the unique, intrinsic, non-linear magnetic response of these SPIONs to the gradient field is directly detected to generate an image.

MPI is the first new imaging modality in 30 years. It was first introduced by Gleich and Weizenecker in 2005 at the Philips Research Laboratory (Germany). Then from 2007, Conolly and Goodwill at the University of California, Berkeley, developed a series of prototype alternative MPI scanners based on the same basic MPI principles but different reconstruction approaches and scanning techniques. Philips later licensed the production of their MPI systems to Bruker BioSpin AG (Switzerland), who subsequently released the world’s first pre-clinical scanner to the market in 2013. Around 2014, Magnetic Insight Inc. (USA) was founded, later becoming the second company to release a commercial pre-clinical MPI scanner in 2016. Other academic labs and companies across the world have also contributed to the development of this technology. Until now, MPI has not been implemented clinically, but multiple groups are working on a human clinical system. Application of MPI has many benefits. SPIONs are a sensitive, safe, and biocompatible tracing material with potentially long physical half-lives, where the MPI signal can remain constant over long time-periods, enabling longer-term imaging of labelled cells. Both the SPION tracers and the scanners themselves have an excellent safety profile, remaining ionising radiation-free. Additionally, MPI is highly applicable in vivo as it provides unambiguous depth-independent detection of SPIONs, with the magnetic field capable of passing transparently through any anatomical tissue or bone without any signal attenuation. Also, without the presence of endogenous background signal, MPI has an essentially infinite contrast, and it can produce highly sensitive and specific images, with detection limits as low as ~200 labelled cells, containing a total of 5.4 ng of iron. Furthermore, MPI is truly linearly quantitative, meaning there is a strong linear relationship between the signal intensity produced and the iron content, and this close relationship holds true for even very small quantities of iron, and at any depth. The coefficient of determination indicates the relationship is almost perfectly linear ($R^2 = 0.99$), thus permitting estimation of SPION concentration in target tissues, based on just the signal...
intensity. Moreover, the temporal resolution of MPI is very high (< 1 second), allowing real-time in vivo imaging and the potential for immediate assistance during medical interventions.49,50

These well-established characteristics make MPI greatly suitable for many applications of clinical relevance, including those previously inaccessible using other imaging modalities. For successful imaging, tailored SPIONs with specific properties are required, as the signal detection in MPI is almost entirely dependent on the properties of the SPION tracer employed. As a consequence, there has been extensive research on the syntheses of novel monodispersed SPIONs, controlling the size, shape, and crystallinity of the core, whether there is any surface modification, and the aggregation state.51-54

Because of the fundamental importance of optimising the characteristics of the SPION for performance in MPI and its applications, the bulk of this review will comprise a comprehensive study of the current research directions in the production of MPI-tailored SPIONs, as well as an in-depth discussion on the up-to-date literature for developments in the current and promising biomedical applications of this rapidly advancing technique. The key areas of focus are divided into applications in theranostics, cell tracking, and perfusion imaging. Prior to this discussion, the simplified principles of image generation in MPI are explained, along with a basic description of the physics and hardware operating within the system. Also included is a detailed comparison between the fundamental properties of MPI and other common imaging modalities with their relative advantages and disadvantages, and a detailing on the recent progress in MPI's clinical translation. Though research in MPI is still in the early stages, we hope this discussion on the major advancements and research directions in this rapidly advancing field over the past 5 years will encourage further exploration into the applications of MPI, as well as in the development of its SPION tracers.

2. MPI background and theory

2.1 Hardware and basic imaging principles

In this section, a simplified description of MPI's hardware and the basic principles for image acquisition and reconstruction is provided. Detailed descriptions of these methods can be found elsewhere.31,46,55,56 To utilise the magnetic properties of SPIONs for imaging, a standard MPI system consists of three major components: a selection field, a drive field, and a receiving coil. In the first MPI scanner developed in 2005,31 the selection field was generated through two permanent magnets positioned such that their field lines are pointing directly towards each other. This alignment produces a strong magnetic field gradient with a sensitive point located in the centre, known as the field free region (FFR), or in this case, the field free point (FFP) (Fig. 1a). The FFP is an area that is void of any magnetic field. This original scanner was able to generate magnetic field gradients of ~3.4 T/m, however, the hardware setup in current preclinical scanners, permits gradients of up to 7 T/m to be reached.57-59

The second component, the drive field, is an alternating magnetic field (AMF). In this original scanner, it is generated by three opposing pairs of drive field coils, one for each respective direction in space (Fig. 1b), each with an amplitude of 10 mT.31 The AMF causes changes in the oscillations of the SPION tracers, which results in variation in their magnetisation that is subsequently detected by the receiving coil. However, the strong magnetic field gradients produced by the selection field saturate all SPIONs except those within the FFP, thus inhibiting the effect of the AMF. Therefore, just SPIONs within the FFP respond to the AMF. Here, it is important to consider the unique magnetic properties of SPIONs in the presence of an external magnetic field.32 First, the two most commonly encountered SPION systems for MPI are single- and multi-core SPIONs, must be described. “Single-core” refers to SPIONs containing just one magnetic core per particle, whereas “multi-core”, containing several closely aggregated magnetic cores per cluster, where the cores are most often linked through dipole-dipole interactions.60,61 This results in their different MPI behaviours, upon exposed to external applied AMFs. Within an MPI system, when SPIONs are exposed to the strong magnetic selection fields in proximity to the magnets, both types of SPION core are fully magnetised to a state of saturation, and therefore, do not generate an MPI signal. In either case, when exposed to small or no magnetic field, as in the FFP, the nanoparticles are randomly oriented and constantly oscillate. The FFP is shifted over the entire field-of-view (FOV) via rapid variation of the drive field. Whenever it crosses a SPION, the magnetic dipole of the nanoparticle flips orientation instantaneously to become aligned with the field lines, and in accordance with Faraday's law of induction, induces a voltage that is detected by the sensitive receiver coils.

This concept can be utilised to generate a 2D projection image of SPION distribution, allowing for spatial encoding of the particle signal, indicating the presence and location of SPIONs within the FOV.62,63 This spatial reconstruction process requires complex algorithms, the most well-established of which, are system function reconstruction (SFR) based processes,31,50,56,64-69 and x-space processes, 32,44,46,55,57,59,70 These two processes rely on the same basic hardware and imaging principles, assigning the signal to a location that corresponds to the respective location of the FFP within the region of interest. The voltages induced in the coil are also linearly proportional to the concentration of SPIONs at the instantaneous location of the FFP, enabling their quantification. This linearity of signal intensity with the density of SPIONs has been confirmed in theory.64 Furthermore, through rotation of the system around the sample, it is possible to capture 2D projection images at multiple angles, where this data can then be converted into a 3D image.

When MPI was first established, SFR-based processes were primarily employed for image reconstruction.31,56,64,65,67,69 These processes require a pre-characterisation of the SPIONs, whose signal response is formulated into a system matrix containing Fourier harmonics of the temporal signal for all possible locations of a point source.52 The system matrix is typically measured physically using a SPION sample,50 but may also be estimated through application of a model.67
Fig. 1 (a) The magnetic gradient field generated by an MPI scanner with two permanent magnets, producing an FFP along with magnetically saturating regions. The FFP is rapidly scanned over the FOV via variation of the drive field, generating signal from the reorientation of the SPIONs' magnetisation. (b) Configuration of drive field coils required to generate a drive field.

Reconstruction of the image is achieved using matrix inversion and regularisation techniques. However, this inversion is often complex since the matrix is large, comprising millions of elements. Additionally, the system matrix is greatly specific to the SPION sample in solution, and thus the accuracy of reconstruction will be less if the SPION behaves differently in tissue, or if the model is inaccurate. Nowadays, x-space processes have received more research attention and are more frequently implemented. In these processes, a fast reconstruction algorithm computes the MPI image without any requirement for pre-characterisation, modelling, and matrix inversions, resulting in robust, real-time imaging. An x-space image is reconstructed from a raw MPI signal through a simple two-step process of velocity compensation of the received signal, followed by gridding of said signal to the instantaneous position of the FFP. Velocity must be compensated in the received signal as the induced signal is proportional to the instantaneous velocity of the FFP.

Recent developments have shown that MPI sensitivity can be significantly improved through application of a more expansive spatial encoding scheme. Instead of scanning the FOV with an FFP, a field free line (FFL) can be utilised. An FFP integrates signal from just a small area, whereas an FFL allows for spatial encoding along a line, integrating signal from an area ~10 times greater. This yields a correspondent theoretical sensitivity increase of a factor of 10, and an increased signal-to-noise ratio (SNR). Additionally, a lower level of power consumption is required. The first experimental setup of an FFL was presented by Knopp et al., where the line is generated by two orthogonal Maxwell coil pairs. The magnetic fields generated by the opposing coils flow in exact opposite directions, thus generating an FFP between each coil pair as before. However, superposition of the generated magnetic fields leads to production of an FFL centered along the bore axis of the scanner. Top et al. recently engineered the first open-sided FFL prototype MPI scanner. Along with the ability to electronically scan the FOV to generate tomographic images, this open-sided design permits interaction with the subject for potential real-time interventional procedures. The results from initial 2D imaging experiments have shown that high quality images with comparatively low resolutions of 2.5 mm can be produced using very low gradient levels (0.6 T/m).

2.2 The Langevin model

In a simplified model, this non-linear magnetisation response of SPIONs in the presence of an AMF follows the classic Langevin magnetisation curve (Fig. 2a), so long as anisotropy, hysteresis effects, and any particle interactions are disregarded. To explain this in terms of an ideal system of single-core SPIONs, or multi-core SPIONs, when the applied field is strong in a particular direction to one side of the particles, the magnetisation starts in a saturated state with the SPIONs aligned in the direction of the field. As the applied field is shifted across the particle, the magnetisation desaturates, eventually to a state where the applied field is ~0 mT, and the SPIONs become randomly oriented. SPIONs have zero remanence and zero coercivity, so as the applied field is further shifted across the particles, there is a linear transition in the curve, shown in Fig. 2a, and the orientation of the nanoparticles flips rapidly with respect to the externally applied AMF. Following this transition, and with a still shifting applied field, the SPIONs re-saturate to the same intensity as before once the field strength has passed a certain threshold, but with reverse polarisation and alignment. The exact opposite process occurs when shifting the applied field in the opposite direction.
it is necessary to first normalise the signal obtained by the PSF of a particular SPION is a measure of the change in gradient strength, it is possible to estimate the overall MPI spatial image resolution. The FWHM is often referred to as the ‘nanoparticle resolution’.81 It by dividing the value for the FWHM of the PSF by the imaging strength, it is possible to estimate the overall MPI spatial image resolution.52,55 It is also possible to compare the FWHM, and thus resolutions, for two different SPIONs. To do so it is necessary to first normalise the signal obtained by the SPION concentration in the sample.

2.3 SPION relaxation

Upon application of the externally applied drive field, the dynamics of SPION magnetisation changes with the relaxation time constant, \( \tau^{-1} = \tau_{\text{Brownian}}^{-1} + \tau_{\text{Néelian}}^{-1} \),2 which is influenced by the Néel time constant (\( \tau_{\text{Néelian}} \)) and the Brownian time constant (\( \tau_{\text{Brownian}} \)). Therefore, it can be stated that the magnetic moments of SPIONs relax to align with the external field through joint Néel and Brownian processes.83-85 The Néel time constant describes the internal flip in magnetisation of the particles from one orientation to another without physical rotation of the particle. This occurs on a timescale of nanoseconds. Whereas the Brownian time constant describes the physical rotation of the particle in space without a change in the internal magnetisation of the particle. This rotation happens on a scale of microseconds. Both time constants are affected by different parameters.83,84 The Néel relaxation process is primarily affected by temperature fluctuation, the particle’s composition and size, effective magnetic anisotropy, and any interdomain interactions within the particles. Conversely, the Brownian relaxation process, whilst also affected by temperature fluctuation and particle size, is influenced by the hydrodynamic volume of the SPIONs, the local microenvironment and the liquid medium viscosity of the immediate surrounding. Though both relaxation mechanisms coexist and often take place simultaneously, in general, SPIONs with smaller core sizes exhibit Néel relaxation to guide the dynamic magnetic responses of SPIONs and produce their signal in MPI, whereas larger core size SPIONs are instead Brownian relaxation dominant.83,85 For further information on these processes, the interested reader should refer to a review written by Krishnan.86

3. Comparison of MPI to other in vivo imaging modalities

Each in vivo imaging technique has its own advantages and disadvantages depending on the application scenario. Table 1 illustrates a basic comparison of the qualities of MPI with other widely applied clinical and pre-clinical modalities. Some of the primary comparative advantages of MPI that can be inferred are its true quantification, high sensitivity and temporal resolution, and radiation-free labelling.

3.1 Nuclear medicine

One of the more popular non-invasive imaging modalities that draws easy comparisons to MPI in terms of its properties is nuclear medicine.87-89 SPECT and PET scans are the two most frequently applied nuclear medicine imaging modalities, with significant clinical potential in that they are highly quantitative, and show great tissue penetration capability.90,91 Both MPI and nuclear medicine are highly sensitive and operate through a ‘hot spot’ detection mechanism of their tracers within a sample. The primary difference between MPI and these techniques, however, is the tracing modality. MPI makes use of SPION tracers, whereas nuclear medicine detects radioactive tracer agents or isotopes. Whilst both MPI and nuclear medicine are highly sensitive with no background signal nor signal attenuation from the tissues, the radionuclides used in PET and SPECT have shorter half-lives on the order of minutes to hours (e.g., PET tracer: \( t_{1/2}\text{(18FDG)} = 2 \text{ h} \); SPECT tracer: \( t_{1/2}\text{(99mTc)} = 6 \text{ h} \)), in comparison to that of SPIONs which has enabled researchers to track the location of SPION-labelled cells for longer time periods (see section 4.2).26 Additionally, SPIONs do not produce harmful ionising radiation as with radionuclides.87,89,92 The shelf life of SPIONs are also orders of magnitude longer, obviating the need for preparation of the tracers immediately before patient use.93 It’s also worth noting that the production costs of SPIONs are significantly lower than those of radionuclides.93

3.2 Magnetic resonance imaging (MRI)

In x-space reconstruction, the imaging effect can be described by a point spread function (PSF).55 A PSF is generated from the differential of this Langevin behavior and provides important information about the signal produced (Fig. 2b). The PSF of a particular SPION is a measure of the change in magnetisation as a function of the applied drive field. There are two important parameters to consider when looking at a PSF: the signal intensity, which reflects the sensitivity of the nanoparticle, and the full-width at half-maximum (FWHM), which is related to the effective spatial dimensions of the signal, and thus the spatial resolution of the nanoparticle. The FWHM is often referred to as the ‘nanoparticle resolution’.81 By dividing the value for the FWHM of the PSF by the imaging gradient strength, it is possible to estimate the overall MPI spatial image resolution.52,55 It is also possible to compare the FWHM, and thus resolutions, for two different SPIONs. To do so it is necessary to first normalise the signal obtained by the SPION concentration in the sample.
Contrast agents/tracer

MRI and MPI is in the physics required for signal generation. MRI utilises a strong field strength, weak gradients, and images across a high uniform magnetic field, whereas MPI utilises a weak field strength, strong gradients and images in the previously described FFR, as generated by the gradient field. Visualising the change in magnetisation via Faraday’s law with a receiver coil in MPI is not dissimilar to the process for image generation in MRI. However, unlike MRI, the magnetisation change in MPI is of electronic, rather than nuclear magnetisation. This contributes to a higher sensitivity in MPI (>2000 ×), as the electronic magnetisation of iron detected in MPI is 22 × 10⁶ times stronger than that of the nuclear magnetisation of water detected in MRI. On another note, where the SPIONs employed by MPI act as a tracing modality detected via ‘hot spots’, the SPIONs that may be employed by MRI act as contrast agents, where the contrast is generated from proton density and the relaxation effects of protons in the vicinity of the particles.

MRI contrast agents are generally differentiated as either “positive” T₁-weighted contrast agents or “negative” T₂ (or T₂*)-weighted contrast agents, where each class manifests proton-spin relaxation times in different ways. T₁-weighted contrast agents shorten longitudinal spin-lattice relaxation times, generating an overall bright image. Conversely, T₂-weighted agents shorten transverse spin-spin relaxation times, generating an overall dark image. SPIONs have many favourable chemical and physical properties that benefit application in MRI, including great magnetic characteristics, targeting capability, limited toxicity, and a unique biodistribution and pharmacokinetic profile. The development of these nanoparticles as MRI contrast agents results in better, safer alternatives to the conventional, toxic, gadolinium-based paramagnetic agents. A number of different parameters including core shape, hydrodynamic diameter, aggregation, and coating choice influence the transverse and longitudinal relaxation times of SPIONs, but the most important determination as to whether SPIONs can be implemented as T₁- or T₂/T₂*-weighted contrast agents is their core size. Generally, larger SPIONs (≥ 5 nm) function as T₂/T₂* contrast agents, whereas smaller SPIONs (≤ 4 nm) function as T₁ contrast agents. The properties of SPIONs ideal for MPI is discussed further in section 4.1. Generally, SPIONs that are used as T₂/T₂* contrast agents can also function as MPI tracers, however they may not be ideal for MRI performance.

SPIONs in MRI are most implemented as T₁ (or T₂*) contrast agents, rather than T₂ contrast agents. Despite them being the most effective T₁ MRI contrast agents to date, with excellent T₂ field-dependent relaxivities surpassing 100 m/M/s, they have several drawbacks. As a negative contrast agent in MRI, SPIONs create ‘black holes’, obscuring the underlying anatomical tissue structures. Additionally, other endogenous sources of contrast may be mistaken for the exogenous SPIONs, such as haemorrhagic tissue or air-tissue interfaces. As a result of this, and since they are not detected directly but instead indirectly, it is not possible to reliably quantify the concentration of SPIONs in targeted tissues. This is in stark contrast to positively-contrasted MPI, which allows efficient quantitation. There have been several studies demonstrating the potential for SPIONs in positively-contrasted T₁ MRI, yet research in this field remains uncommon. In a recent paper, Than et al. synthesised monodisperse SPIONs, with sizes ≤ 5 nm, through application of a millifluidic multistage flow reactor. These flow-synthesised SPIONs generated great values for enhancement of the T₁ contrast, with longitudinal relaxivities (r₁) greater than 10 mM⁻¹ s⁻¹, and transversal relaxivities (r₂) reduced to just 20.5 mM⁻¹ s⁻¹. On a final note, scanning and imaging in MPI is much more rapid and straightforward than in MRI, with no specialised training required to acquire or interpret the images.

3.3 Multi-modal imaging with MPI
A key disadvantage of modalities like MPI and nuclear medicine is that the images produced do not provide any anatomical information. They are only able to generate morphological information from contrasted structures. Therefore, to provide context about where the SPIONs have accumulated in the animal, MPI requires an anatomical reference. MRI and CT are examples of anatomical imaging modalities. They may provide complementary images of the structural information of a sample, with which an MPI signal can be referred against.

Dual modality MPI systems are currently being investigated, looking to combine the advantages of two techniques within one system, similar to how PET/MRI or PET/CT systems complement each other.\textsuperscript{122-124} MPI is most frequently co-registered with MRI as it is possible to employ the same type of SPION for both techniques, and because both modalities utilise magnetic fields to generate their images. Typically, the imaging is performed in their respective devices and the information is combined via post-processing procedures to produce a 3D tomographic image of the sample.\textsuperscript{125,126} The first in vivo studies for co-registered MRI/MPI images were conducted by Kaul et al., in which measurements with preclinical 7 T MRI were performed before and after MPI scan.\textsuperscript{125}

Recently there has been lots of work on the construction of hybrid MPI/MRI scanners to help ease the co-registration process.\textsuperscript{123,124} High spatial resolution accuracy can be achieved as MRI and MPI modalities share the same FOV. However, both modalities require very different magnetic field topologies, and the field strength required for MRI is so great that the SPION tracers become fully saturated. Consequently, simultaneous imaging is difficult to realise, and acquisition of data sequentially has been employed instead. The first instrument setup to combine both modalities within a single system was introduced by Franke et al. in 2013, wherein, all magnetic components are arranged concentrically and with an identical magnetic centre.\textsuperscript{127} The subject can therefore be sequentially measured with both modalities without the need for repositioning or transportation of the subject. From this initial magnet design, the first fully integrated pre-clinical MPI/MRI system for static 3D imaging was presented.\textsuperscript{123,128} Successful initial phantom measurements were taken, demonstrating the feasibility of this system. The first in vivo results from this new methodology were gathered later by the same group,\textsuperscript{129} working through the integration of Halbach rings into a rotating gantry.\textsuperscript{72} This offers an open design, providing a ‘window’ for direct feedthrough of X-rays and thus, CT imaging. As a result, the quantitative SPION distribution as well as the anatomical information of the surrounding tissue material can be visualised simultaneously and rapidly, providing a potential basis for improving diagnostic accuracy in pre-clinical imaging.\textsuperscript{133}

### 4. Development of SPIONs as MPI tracers

#### 4.1 Introduction to SPIONs

Signal detection and imaging quality in MPI is almost entirely dependent on the specific SPION tracer used.\textsuperscript{53,134} This stems from direct detection of the non-linear magnetisation of SPIONs by the system for signal generation, without any background interference from tissues or signal noise. Hence, for improved MPI performance, tailored tracers with specific properties are required. Generally, SPIONs used in MPI comprise a spherical crystalline core, typically of maghemite ($\text{Fe}_3\text{O}_4$) or magnetite ($\text{Fe}_3\text{O}_4$) crystals, which are colloidal stabilised using biocompatible magnetically neutral polymeric coatings like polyethylene glycol (PEG) or carboxydratex. These particles tend to have a hydrodynamic diameter between 50 and 100 nm.\textsuperscript{135} At this size, iron oxide nanoparticles show superparamagnetic behaviour, with zero remanent magnetisation and coercivity following removal of the field, in the case of multi-core SPIONs or a system of single-core SPIONs.\textsuperscript{136,137} This intrinsic property of SPIONs, together with the non-linear Langevin behaviour detailed above,\textsuperscript{80} allows for signal differentiation and detection of SPIONs in MPI.

The application of SPIONs as a tracing material is highly advantageous, not just in MPI, but in other SPION-compatible imaging modalities also. This was demonstrated in section 3.2, through a discussion of how MRI performance can be enhanced through implementation of SPION contrast agents. Generally, SPIONs are widely available, easy to handle, and relatively inexpensive compared to other commonly used tracers.\textsuperscript{82} Furthermore, SPIONs are non-radioactive, and their signal does not decay over time. This enables effective longitudinal tracking studies of cell-based therapeutics.\textsuperscript{26} Outside of MPI, SPIONs have been implemented in a wide variety of applications. They have shown promising clinical indication in iron supplementation therapy for anaemic candidates,\textsuperscript{138,139} cell separation,\textsuperscript{140,141} drug delivery,\textsuperscript{142} hyperthermia,\textsuperscript{143} mapping of lymph node metastases,\textsuperscript{144,145} diagnosis of liver cancers,\textsuperscript{146,147} and as a $T_1$ agent for angiographic MRI,\textsuperscript{148} to name a few. There are several SPIONs that have either received approval from the USA Food & Drug Administration (FDA) for clinical applications or are in/close to a clinical trial.\textsuperscript{149,154} A selection of these formulations, as well as some well-established pre-clinical SPIONs that could potentially serve as tracers in MPI, are presented in Table 2.

A particularly important feature of SPIONs, which is true of all nanomedicine, is that there is great modularity and flexibility possible when altering the structure and properties of the nanoparticle. This tunability is particularly valuable in MPI applications due to the wide range of properties that can be engineered into these materials. The following section provides a discussion on the synthesis and application of SPIONs, which is a key focus of the current literature in the field.
where the advantages of a particular SPION in a particular application is dependent on so many factors, including the size and shape of the iron oxide core, the type of surface coating, and whether there is any additional surface modification. As a result, researchers can produce meticulously engineered tracers in MPI with different physical and biological properties specific for different applications (Fig. 3).

**Table 2** Well-documented examples of clinical or pre-clinical SPION formulations that have been used or may have potential as MPI tracers.

| Name                     | Company                          | Coating               | Hydrodynamic diameter (nm) | Core diameter (nm) | Applications                        | Market status          |
|--------------------------|----------------------------------|-----------------------|---------------------------|-------------------|-------------------------------------|------------------------|
| Ferucarbotran\(^{22}\)   | Bayer AG (Resovist*/VivoTrax*)   | Carboxy dextran       | 62                        | Multi-core, *4 each core | MRI contrast agent & MPI             | Clinically approved (Resovist* in EU, Japan) |
| Resovist*/VivoTrax*      | Magnetic Insight (VivoTrax\(^{TM}\)) |                       |                           |                   | Iron supplementation therapy & MRI angiography | Clinically approved (USA) |
| Ferumoxytol\(^{152}\)   | AMAG Pharmaceuticals (Feraheme*), Takeda Pharma (Rienso*) | Carboxymethyl-dextran | 30                        | 3-4               | MRI blood pool agent                | Clinical trial         |
| Feronio* (EU)           |                                   |                       |                           |                   |                                     |                        |
| Feruglide\(^{152}\)     | GE Healthcare                     | PEGylated starch      | 20                        | 5-7               | MRI blood pool agent                | Pre-clinical           |
| Clariscan™              |                                   |                       |                           |                   |                                     |                        |
| FeraSpin* XXL\(^{158}\) | Millen Biotec                     | Carboxy dextran       | 65                        | Multi-core, 5-7 each core | MRI blood pool agent                | Pre-clinical           |
| LS-008\(^{159}\)       | LodeSpin Laboratories MicroMod    | PMAO*-PEG             | 80                        | 25                | MRI blood pool agent                | Pre-clinical           |
| Perimag\(^{159}\)      |                                   | Dextran               | 130                       | Multi-core, *5.5 each core | MRI contrast agent & MPI             | Pre-clinical           |
| PrecisionMRX\(^{150}\)  | Imagion Biosystems               | mPEG                  | 41                        | 24-25             | Hyperthermia & MRI                 | Pre-clinical           |
| Synomag*-D\(^{151}\)   | MicroMod                          | Dextran               | 56                        | Multi-core, 5-15 each core |                                     |                        |

*poly(maleic anhydride alt-1-octadecene).

At the beginning of MPI exploration, SPION tracers that were designed as \(T_1\) or \(T_2\)-weighted MRI contrast agents were evaluated for their MPI performance. However, since MRI and MPI have totally different physics, it was determined that these existing tracers were not ideal for reliable application in MPI and that new SPIONs would have to be tailored to MPI’s unique set-up.\(^{3,5,63,162}\)

Multi-core SPIONs are far more common than their single-core counterparts.\(^{5,158,163,164}\) Many of the frequently implemented commercial SPIONs have multi-core structures, as demonstrated in Table 2. It is, however, very complex to control the structural uniformity of clustered cores, as there are many structural variables that need to be considered, including, the number of cores per cluster, the cluster diameter, shape, density, and the inter-core distances and spatial distribution.\(^{50}\) Any alteration in these parameters will lead to considerable changes in the packing arrangements, and this can significantly affect the dipole-dipole or exchange-interactions between the tightly associated cores, which strongly influence the magnetic behaviour of the SPIONs.\(^{165-167}\) Because of this, in general, multi-core SPIONs have less uniform magnetic properties than standard single-core formulations, and produce poorer signals in MPI. Despite this, the presence of multiple cores can also be advantageous, if there is proper control of the structure during the synthesis.

Many recent studies on the synthesis of single-core SPIONs for MPI have been focussed on the effect of core size on MPI performance and sensitivity. One of the principal reasons for the poor performance of MRI-tailed SPIONs in MPI is the small size of their magnetic iron oxide cores (generally < 10 nm), which results in lower magnetic moments for the nanoparticles.\(^{53,168}\)

Both core structures hold equally important roles in MPI applications. Individually coated single-core SPIONs have demonstrated significantly increased blood half-lives *in vivo*,\(^{169}\) favouring their use in perfusion imaging.\(^{58,170,171}\) Additionally, the structural uniformity is beneficial for the targeted delivery of SPIONs to a therapeutic site, since core size is a major determinant of a SPIONs pharmacokinetic behaviour, and therefore its biodistribution.\(^{172}\) Multi-core SPIONS have also been implicated as greatly beneficial for many biomedical applications. On the other hand, the presence of the magnetic coupling interactions between the clustered cores is particularly advantageous in magnetic hyperthermia applications.\(^{165}\) The improved magnetic moment demonstrates advantages in standard magnetic hyperthermia techniques,\(^{173-177}\) and MPI-
coupled magnetic fluid hyperthermia (MFH), otherwise called MPI-MFH.\textsuperscript{178-181} The benefits of MPI-MFH is discussed in depth in section 5.2.2.

The shape of the core, or cores in the case of a multi-core formulation, is another factor that demonstrates strong influence over SPION performance in MPI. To date, most SPION cores developed for MPI, and its applications, have a spherical shape. However, it is known that non-spherical MNPs can offer significant advantages in many different biomedical applications, as a consequence of altered physical properties, and the potential for improved magnetic characteristics, including Ms, magnetic anisotropy,\textsuperscript{182-184} and heating properties.\textsuperscript{185,186} Additionally, as alternative MNP shapes present larger available surface areas for cell interaction, as compared with equivalently sized spherical nanoparticles, they often demonstrate greater cellular uptake.\textsuperscript{187} As well as this, they have potential for enhanced blood circulation half-lives.\textsuperscript{188}

In theory, the potential for improved magnetic properties, through the synthesis of non-spherical SPIONs, should be an effective way enhance MPI sensitivity and spatial resolution. As a result, such work has attracted the attention of various research groups. In particular, there has been sustained recent interest in the implementation of cubic SPIONs in MPI.\textsuperscript{189,190} These nanocubes have a lower proportion of disordered spins at their surface and smaller surface anisotropies in comparison to equivalently sized spherical SPIONs.\textsuperscript{190} This results in higher values for Ms and magnetic susceptibility, and consequently improved MPI performance, at certain sizes.\textsuperscript{191} Along with this enhanced magnetic performance, the tendency of cubic SPIONs to spontaneously form chain-like arrangements has led to improvements in the performance of SPIONs in MPI-MFH.\textsuperscript{13,191,192} A recent study from Avugadda et al. showcased the advantages of SPIONs, comprising a controlled number of cubic cores, on MPI, and potential MPI-MFH performance.\textsuperscript{5} Among the variety of magnetic assemblies synthesised in this study (Fig. 4a), multi-core dimer and trimer structures exhibited the greatest MPI properties (Fig. 4b and c). The other structures investigated were larger multi-core clusters of nanocubes, and individual single-core nanocubes, all synthesised using the same polymer coating. The enhanced performance is attributed to the beneficial uniaxial magnetic dipolar coupling present in the chain-like smaller multi-core assemblies.\textsuperscript{193}

Another key consideration in the design of SPIONs for MPI applications is the choice of biocompatible surface coating. As a general trend, a coating must confer colloidal stability to the particle, assisting aggregation prevention under various complex physiological environments, whilst also mediating any interactions with biological entities. This further stability can also potentially prolong half-life in the bloodstream, help prevent agglomeration during storage or application, and counteract possible oxidation.\textsuperscript{108,109} It is also important to ensure the coatings promote cellular uptake yet preserve the optimal magnetic response of the SPIONs when dispersed in the acidic endosomal environment. There are many common choices of coating, each with their own advantages and functions depending on their molecular structure. Polymers are among the most popular coatings for SPIONs.\textsuperscript{195,196} Early MPI research typically relied on multi-core ferucarbotran (Resovist, Bayer AG), a SPION originally clinically approved as an MRI contrast agent in the liver. Ferucarbotran is coated by a carboxydextran polymer.\textsuperscript{50} More recently, PEG and dextran polymer coatings have been employed most frequently. This is because they are not readily recognised by macrophages in the liver and spleen when administered intravenously and consequently have enhanced circulation time.\textsuperscript{107,108} They have also been generally recognised as safe by the FDA.\textsuperscript{108} The biodistribution of PEG- and carboxydextran-coated SPIONs was studied by Keselman et al.\textsuperscript{199} While the carboxydextran-modified SPIONs are cleared rapidly to the liver, the PEG-coated particles are sustained for a relatively long blood half-life of 4.2 hours before eventual excretion through the RES system, illustrating the benefit of PEG-coated SPIONs for \textit{in vivo} studies.

It is also worth noting that coatings can be altered to tailor their biochemical properties towards a specific physiological application. Examples of such alteration are that the coating may be grafted differently (\textit{i.e.}, it can be chemisorbed, physisorbed, covalently bonded \textit{etc.}), or that different graft densities and molecular weights of coating may be applied. Guzy et al. demonstrated the importance of polymer coating choice, and coating molecular weight, on SPIONs undergoing biodegradation.\textsuperscript{81} SPIONs can undergo a variety of physicochemical changes as they degrade, which generally results in detrimental effects on MPI signal properties and ‘nanoparticle resolution’. It was found that larger polymers with a greater molecular weight will degrade more slowly in harsher

![Fig. 4 Magnetic assemblies comprising a controlled number of cubic cores, designed for optimal MPI performance. (a) Transmission electron microscopy (TEM) images of the different SPION assemblies made from iron oxide nanocubes encapsulated in a poly(styrene-co-maleic anhydride) coating. The assemblies synthesised were single nanocubes (Si), short-chain dimers/trimers (Di/Tri), and various 3D-Cluster configurations. (b) PSF demonstrating the MPI signal obtained for each synthesised sample, in comparison to that of the reference, VivoTrax (Std). (c) Histograms of the corresponding SNRs for each synthesised sample and the commercial reference.](https://creativecommons.org/licenses/by/4.0/) Copyright 2020, the authors. Article reproduced from Ref. 5 with permission from the Multidisciplinary Digital Publishing Institute. This is an open access article distributed under the terms of the CC BY License (https://creativecommons.org/licenses/by/4.0/).
endosomal conditions, such as at a tumor site, and thus their MPI signal will remain for longer.

The surface coating can also act as a structural support for additional surface modification, with the potential to conjugate a huge variety of possible functional molecules like drugs and ligands to improve molecular targeting, and proteins, antibodies, or aptamers for highly specific chemical interactions with complex biological systems. It can also provide a platform for imaging tags, specifically molecules that allow dual modality imaging with MPI, for example fluorophores for fluorescent microscopy.\textsuperscript{4,200} Following successful surface-modification, the resulting SPIONs may be employed for MPI.

4.2 Synthetic methods

The modularity in SPION design in Section 4.1 is incredibly useful. To ensure production of SPIONs with desired properties, it is fundamentally important to choose the most appropriate method of synthesis. The choice has a strong influence on key physical characteristics of SPIONs such as crystal structure, core size, and size distribution, and consequently, on the magnetic properties of the nanoparticles. A comparison between five of the most employed synthetic methods are briefly summarised in Table 3, demonstrating their specific advantages and downsides. Besides those outlined in the table, SPIONs can be effectively prepared by various other techniques, including sonochemical and electrochemical deposition, microwave irradiation, laser pyrolysis, and reduction methods.\textsuperscript{195,201-203}

Because of their mostly beneficial properties, thermal decomposition and co-precipitation are preferred for the synthesis of SPIONs for MPI.\textsuperscript{204} Co-precipitation works by the simultaneous precipitation of Fe\textsuperscript{3+} and Fe\textsuperscript{2+} aqueous salts following addition of a basic solution.\textsuperscript{206} It is a cost-effective and simple process, capable of high-yielding, scalable syntheses. Additionally, the synthesis does not require an organic solvent and the precursors used are generally environmentally friendly.\textsuperscript{206} Despite these beneficial qualities, the nanoparticles formed are often relatively polydisperse with a low degree of crystallinity.\textsuperscript{207,208} This can result in an appreciably weakened MPI signal. Differently, SPIONs with a very narrow size distribution and excellent crystallinity can be prepared through thermal decomposition.\textsuperscript{209,210} In this method, SPIONs are synthesised through the decomposition of organoiron precursors in organic solvents with high-boiling points, in the presence of stabilising surfactants. These surfactants bind to the growing nanocrystals, controlling their nucleation and growth. However, despite formation of high-quality samples, this technique requires expensive, and generally toxic reagents and solvents, and is therefore not environmentally friendly.\textsuperscript{208} Furthermore, as a hydrophobic coating is formed on the surface of the SPIONs during synthesis, an additional surface modification step is required to obtain the biocompatible, water dispersible SPIONs to be used in biomedical applications. An in-depth description of all syntheses techniques is not within the scope of this review, thus, for further description of the other techniques, interested readers should refer to reviews by Thanh et al.\textsuperscript{211,212}

4.3 Recent developments in SPION research

Similar to when SPION contrast agents were first realised for MRI in the 1980s,\textsuperscript{211} research on the development and synthesis of monodispersed novel MPI-tailored SPIONs has recently become an important area of research.\textsuperscript{5,6,43,189,214-217} These new particles can be functionalised and optimised for improved performance in specific applications, like for increased circulation time or for more efficient cell targeting. Whilst MPI-optimised tracers will have to undergo a lengthy evaluation before clinical approval, the exploration and development of better performing SPIONs should spur further work towards clinical applications.

The development of tracers with long circulation times is crucial for many applications. Typically, when nanoparticles are administered into bloodstream circulation, there is just a narrow time window to image the particles before they accumulate in the liver and spleen for excretion, where the concentration of the particles falls such that MPI can detect no meaningful signal.\textsuperscript{103} Studies on the circulation time of carboxydextran-coated multi-core ferucarbotran (Resovist), which is generally considered the ‘gold-standard’ of MPI tracers, show that following administration to rabbits, the MPI signal decreased to ~12% of the initial intensity after just 15 min, and within 30 min, the signal had disappeared entirely.\textsuperscript{218} Another commercially available tracer with potential use in MPI is dextran-coated Synomag-D (MicroMod), where the MPI performance shows better circulation times (t\textsubscript{1/2}, ~1 h) as compared to Resovist.\textsuperscript{219} Khandhar et al. developed a new single-core nanoparticle known as LS-008 (LodeSpin Lab) through a post-synthesis oxidation method.\textsuperscript{210} This MPI-tailored tracer of core diameter, 25 nm, produced outstanding resolutions (1.6 mm at a 7 T/m/μm field gradient) and was designed for long blood circulation times (t\textsubscript{1/2}, ~105 min in mice), with an exceptionally stable PMAO-PEG coating. However, for many of the potential applications of MPI, much longer half-lives are required.\textsuperscript{3,5,8,9,170,171,220} Additionally, the availability of longer-circulating tracers will reduce the quantity of tracer required and/or the number of tracer administrations for treatment.

New MPI tracers with long blood circulation half-lives of 7 hours were obtained by Liu et al., termed RL-1, as shown in Fig. 5a.\textsuperscript{20} These single-core SPIONs were developed using a semi-batch thermal decomposition process with molecular oxygen addition. This was followed with an optimised PEG-silane ligand exchange process, producing SPIONs with high values for spatial resolution (~2 mm at 5.7 T/m) and sensitivities greater than multi-core Synomag-D, and ~3 times greater than multi-core ferucarbotran (Resovist) (Fig. 5b). Another long-circulating tracer was developed by Song et al.,\textsuperscript{20} composed of a Janus iron oxide @ semiconducting polymer nanostructure (Fig. 5c) synthesised through a nanoprecipitation method reported prior by the same group, shown in Fig. 5d, where the semiconducting polymers employed demonstrate great biocompatibility.\textsuperscript{12} These particles provide a nanoplatform for ultrasensitive multi-modal imaging (MPI/MRI/photoacoustic/fluorescence), termed MMPF nanoparticles) of tumor xenografts in living mice,
An alternative approach to increasing circulation times includes entrapping the SPIONs into human red blood cells (RBCs). Through nuclear magnetic resonance measurements, such loaded RBCs were demonstrated to circulate for over 12 days in mouse models before an obvious reduction in concentration could be detected. For RBCs loaded with Perimag, transmission electron microscopy images show that the particles have a spatially uniform distribution within the cells, without any discernible indication of particle aggregation. Utilising Resovist-loaded RBCs, Rahmer et al. presented the first evidence that SPION-loaded cells could be imaged in vivo with MPI, showing clear imaging of the blood pool in mice several hours following injection. This observation was supported with magnetic particle spectroscopy measurements, performed to determine the concentration of iron in samples of blood extracted from the mice at different time points following injection. Antonelli et al. also performed a study encapsulating different commercially available SPIONs, Synomag-D and Perimag (MicroMod) into RBCs through hypotonic dialysis. COOH-functionalised Perimag loaded RBCs proved to be viable cells, while the magnetic signal of the equivalently functionalised Synomag-D loaded cells dropped sharply. Therefore, just the Perimag-loaded RBCs have potential for MPI diagnostic applications, showing potential for longer blood retention times than the equivalent free nanoparticles. Successful application of MPI to the imaging of pathological diseases depends on the quantity of nanoparticles that accumulate at a diseased site relative to other sites. Hence, another important area for MPI-tailored tracer development is in the synthesis of nanoparticles with coatings functionalised towards the active targeting of specific pathophysologies.

One of the more prevalent functional targeting applications is the targeting of nanoparticles towards cancerous cells/tumors. Cancer is one of the global leading causes of death, accounting for almost 10 million deaths worldwide in 2020 alone. Interest in its effective targeting is therefore necessary. Arami et al. conjugated the glioma-targeting glycoprotein, lactoferrin, to the PMAO-PEG surface coatings of their optimised single-core MPI tracers with diameters of 25-27 nm. Very high-resolution 3D tomographic multi-modal images permits the tracer to be tracked and quantified longitudinally for up to 85 days (Fig. 5e). DOI: 10.1039/D1NR05670K

| Synthetic method   | Synthesis complexity | Reaction temperature (°C) | Reaction length | Solvent       | Size distribution | Shape control | Yield          |
|--------------------|----------------------|---------------------------|-----------------|---------------|------------------|--------------|----------------|
| Co-precipitation   | Very simple, ambient conditions | 20-90                      | Minutes         | Water         | Relatively narrow | Poor          | High, scalable |
| Thermal decomposition | Very complicated, inert atmosphere | 100-320                   | Hours-days      | Organic compound | Very narrow     | Very good    | High, scalable |
| Hydrothermal       | Simple, high pressure  | 150-220                    | Hours-days      | Water-ethanol | Relatively narrow | Very Good    | Medium, scalable |
| Microemulsion      | Complicated, ambient conditions | 20-50                      | Hours           | Organic compound | Relatively broad | Good         | Low, not scalable |
| Sol-gel            | Complicated, ambient conditions | 25-200                     | Hours           | Water-ethanol | Relatively broad | Good         | Medium, scalable |

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![Fig. 5](https://example.com/fig5.png)
(MPI/CT/X-ray) demonstrate an enhanced uptake of the functionalised SPIONs in brain cancer xenografts in mice (Fig. 6a and b). This is due to the fact that lactoferrin molecules can pass through the blood brain barrier (BBB) with ease, through a receptor-mediated transcytosis mechanism, for active targeting. Another work in this field, by Tomitaka et al., investigated the functionalisation of gold-coated multi-core SPIONs of ~30 nm with transferrin (Fig. 6c). Like lactoferrin, transferrin is a brain gloma targeting ligand, which has exhibited great specificity due to the high expression of transferrin receptors on the surface of brain gloma and capillary endothelial cells. They demonstrated a high functionalisation efficiency of 58% for the SPION with the targeting ligand, using the procedure in Fig. 6d, and these functionalised particles showed great biocompatibility also.

Another targeting application, which is receiving increased interest, is the targeting of active myeloperoxidase (MPO), a potential inflammatory marker of vulnerable atherosclerotic plaque. Tong et al. developed novel single-core multi-modal SPIONs, as termed 5FeC nanoparticles conjugated with 5-hydroxytryptamine and a cyanine 7 N-hydroxysuccinimide ester, that can specifically target MPO and identify these high-risk plaques in vivo. These 21 nm nanoprobes image the active MPO in an atherosclerosis mouse model with high sensitivity, thus enabling quantitative evaluation of the severity of inflammation and monitoring of the MPO activity.

4.4 Effect of SPIONs on spatial resolution and sensitivity in MPI

Traditionally, the most significant technical weakness of MPI is the relatively poor spatial resolution (typically ~1 mm). To compete with preclinical CT and MRI, improving these resolutions to the submillimetre range would be an enabling advance. MPI spatial resolution can be defined as the ability to clearly distinguish two signals with the same intensity in space. From a physical and radiologic viewpoint, this depends on two major factors. First is the magnetic gradient strength of the instrument, more specifically, the stronger the gradient, the narrower the FFR, allowing assignment of the generated SPION signal to a narrower space, and thus a greater spatial resolution. The second factor comes from the physical and magnetic properties of the SPION utilised. This is based on effects to the FWHM of the PSF, where nanoparticles with narrower PSFs produce higher achievable resolutions. Along with a higher quality image, utilising a SPION with better spatial resolutions could significantly reduce instrument cost. A 10-fold improvement in resolution from MPI-tailored SPIONs could potentially reduce the cost of a clinical instrument by up to 100-fold, as spatial resolution can be exchanged for lower MPI gradients which would be required to be high in a clinical system.

Sensitivity is another important parameter in determining overall MPI performance. A high sensitivity permits the detection of very small amounts of tracer, and this is enabling for many biomedical applications, and is particularly important in cell tracking. It is characterised by the height of the Langevin curve at the transition point, where a taller curve indicates a higher sensitivity. Consequently, a greater signal intensity in a PSF, also signifies a better sensitivity. As with spatial resolution, progressing to the theoretical limits of MPI for high sensitivity involves work on both the hardware and tracers. In terms of instrumentation, advancements in sensitivity have most frequently been from developments in coil design. Regarding tracer development for enhanced sensitivity, the signal intensity is governed by the physical and inherent magnetic properties of the tracer, as with spatial resolution, but is mostly dependent on the Ms, and as a result, hampers performance. An empirical ideal curve at the transition point, where a taller curve indicates a higher sensitivity.

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Given what has been discussed, the development of SPIONs with optimal sensitivity and spatial resolution performance has become a crucial part of MPI research. Sustained work in this area will enable high-performing, cost-effective, and safe human MPI. Tailoring of the structural parameters of SPIONs, such as their shape and crystallinity, can increase resolution and signal strength through enhancing of Ms. One of the most important factors for improved performance is the size of the magnetic core. For single-core SPIONs, resolution and sensitivity increase cubically with core size, stemming primarily from higher Ms values. Unfortunately, Tay et al. demonstrated that the improvement in performance is limited with respect to size, as a result of increased Brownian relaxation blurring and a gradual shift from the superparamagnetic to ferromagnetic regime as sizes increase. This has been described as the Langevin wall, and this phenomena reduces Ms, and as a result, hampers performance. An empirical ideal core size with respect to both resolution and sensitivity performance has been estimated at 24-28 nm for single-core SPIONs. Gevaert et al. examined the MPI performance of several commercially available SPION tracers, confirming the resolution and sensitivity dependence of SPIONs to core size.

Recently, a new strategy for tracer formation has been established based on strongly interacting particle-particle interactions. This has been achieved through the development of SPIONs with functionalised surfaces, allowing for efficient targeting towards specific cell populations. These strategies have demonstrated enhanced uptake and imaging capabilities in various in vivo models, offering promising avenues for future applications in both preclinical and clinical settings. Further advancements in this field are likely to lead to improved diagnostic tools and therapeutic approaches, ultimately extending the potential of MPI as a versatile imaging modality.
magnetic dipole interactions which subsequently form nanoparticle chains in fluid, on application of an external magnetic field (Fig. 7a). This chaining was found to amplify the applied field 10-fold, resulting in very sharp PSFs. Through this method, Tay et al. demonstrated the potential of a 40-fold boost in sensitivity, and unprecedented 10-fold improvements in spatial resolution (Fig. 7b). These properties theoretically allow the tracking of single cells in vivo. There have been several other studies testing the ‘chaining hypothesis’, specifically, examining the parameters that effect chain formation times. Colson et al. outlined that for optimised chain formation time, the conditions required are low media viscosities, as viscous solvents can block chain formation, and high nanoparticle concentrations, as the smaller the inter-particle separations, the easier it is for dipoles to interact.

The formation of these ‘chained’ tracers has also been investigated and modelled computationally. Zhao et al. employed simulations to evaluate the effect of the magnetic dipole-dipole interactions on the MPI performance and dynamic magnetisation of individual particles within the chain. The results illustrate similar MPI signal intensity and resolution enhancements to those demonstrated by Tay et al., for interacting chains of 2 particles. They also suggest a large parameter space for design that can be used to tailor these chains towards optimised MPI performance, including, the number of particles in the chain and their separation distances, the composition of the SPIONs, and the viscosity of the solution.

Outside of novel ‘chaining’ approaches, there are other more established methods to improve spatial resolution and sensitivity through tracer synthesis, one of which is through the synthesis of SPIONs with improved monodispersity. Monodisperse SPIONs with uniform magnetic domains have smaller FWHMs and higher values of Ms in the PSF, resulting in nanoparticles with higher spatial resolutions and signal intensities. Dadfar et al. describe a straightforward precipitation synthesis and following sequential centrifugation protocol to obtain monodisperse single-core particles of (polydispersity index (PDI) below 0.1), these optimised dispersions showed substantially improved Ms values, and thus performance in MPI, MRI and MFH, up to 7 times greater in comparison to the polydisperse starting formulation, as well as to commercially recognised SPIONs, such as Resovist. However, this is a two-step process and is not feasible for scale-up, nor is environmentally friendly. In a different work, Unni et al. synthesised monodisperse SPIONs through a modified thermal decomposition process involving the controlled addition of molecular oxygen. The particles synthesised in oxygen presence demonstrate greater magnetic properties (Fig. 8a), and exhibit a better MPI performance (Fig. 8b) than those in the absence of oxygen, with greater Ms values (74 Am²/kg vs 17 Am²/kg, respectively), and consequently improved values for FWHM (12.0 mT vs 15.2 mT, respectively) and signal intensity (39.0 mL/mg(Fe) vs 13.1 mL/mg(Fe), respectively). This improvement is attributed to appreciably more uniform magnetic and physical domain sizes, and fewer structural defects.

The choice of coating should also not be neglected when optimising SPIONs for higher image resolutions and sensitivities. In one study, Horvat et al. reported the positive impact of preparing SPIONs with cross-linked over non-crosslinked polymeric coatings on performance in MPI. The cross-linked single-core SPIONs, of average core size 18 nm, termed PF127DAPG, displayed a superior SNR, and consequently higher spatial resolution than their uncross-linked equivalents, PF127. Resulting from this, an iron quantity of 5.3 μg was required in PF127 SPIONs to produce the same spatial resolution values as 1.3 μg of iron in PF127DAPG SPIONs.

4.5 Development of alternatives to SPIONs as MPI tracers
 Whilst almost all tracers designed for MPI have been based on iron oxide species, there has, in recent years, been evidence to suggest the potential of alternative MNPs to SPIONs for MPI, however research in this area remains underdeveloped. Theoretically, any MNP that is both highly biocompatible, and can display superparamagnetic behaviour with favourable Ms values, could potentially be employed as an effective MPI contrast agent.

Iron nanoparticles have generated exceptionally high values for Ms, up to 176 emu g$^{-1}$, at sizes of 13 nm. However, as with many metal MNPs, they suffer from poor stability and this has limited their application in imaging. Such nanoparticles are rapidly oxidised to iron oxide following exposure to air, and therefore require stabilisation for application in MPI. Gloag et al. prepared superparamagnetic single-core nanoparticles with zero valent iron cores, coated with an iron oxide shell and a strongly binding brush co-polymer (Fig. 9a and b). The iron oxide coating prevents the rapid oxidation of the metallic core, and the additional polymeric layer provides high colloidal stability, where the nanoparticles could remain water-dispersed for over 8 weeks. These nanoparticles show an excellent Ms of 166 emu g$^{-1}$, at a size of 14 nm (Fig. 9c). At this size, the coated nanoparticles achieve an MPI signal intensity that is ~80% that of the much larger multi-core VivoTrax (Fig. 9d), whilst also having a very similar spatial resolution. In comparison, SPIONs of a similar size could not generate recognisable signals in MPI, due to their weak values for Ms. Strong MPI properties for tracers at this size is significant, opening the door to many potential MPI applications within cells and the brain, that were not possible with larger nanoparticles.

Metal alloy MNPs have also become of interest for MPI and its applications. FeCo alloyed nanoparticles in particular, demonstrate almost unmatched Ms values up to 215 emu g$^{-1}$. On top of this, they exhibit superparamagnetic behaviour at sizes less than 20 nm, indicating their potential for MPI. However, as with iron nanoparticles, they must be stabilised to prevent oxidation on exposure to air. In a recent study, Song et al. synthesised 10 nm FeCo nanoparticles (Fig. 10a) coated with a layer of graphitic carbon and PEG. These exhibit exceptional MPI signal intensities, 6.08 times higher than VivoTrax for equivalent molar core concentrations (Fig. 10b). The graphitic carbon coating prevents rapid oxidation of the unstable FeCo core. Many more alloyed nanoparticles exhibit promising properties for MPI application, including FePt (Ms = 100 emu g$^{-1}$) and Fe$_3$C (Ms = 125 emu g$^{-1}$) particles, but none, as of now, have been investigated for MPI.

Ferrites of the Mn$_x$Fe$_{3-x}$O$_4$ general formula, display a spinel structure where M could theoretically be any divalent transition metal ion. In the standard Fe$_3$O$_4$ SPION structure, M is Fe$^{2+}$. Doping of this structure with appropriate metal ions can improve the MPI performance of the particles by altering their magnetic properties. Upon doping, the Fe$^{2+}$ ion is substituted to a desired extent with the new divalent cation dopant (e.g., Zn$^{2+}$, Co$^{2+}$, or Mn$^{2+}$). However, for potential use in biomedical applications, biocompatibility and toxicity of the ferrites is a concern.

In one recent study, Silvestri et al. synthesised cubic ferrite nanoparticles with a tunable quantity of doped Co and Zn, which were then analysed for their MPI capability. Firstly, they synthesised high quality Co-ferrite through a non-hydrolytic synthesis procedure (Fig. 11a). By altering the metal precursors used in this process, the cobalt ions could either be partially substituted with zinc ions, producing mixed Zn-Co-ferrite (Fig. 11b), or totally substituted, producing Zn-ferrite (Fig. 11c). All the cubes were synthesised within a similar size range below 15 nm.
One of the earliest applications in MPI was the systematic tracking of cells, as with MRI.\textsuperscript{257-259} MPI benefits in vivo cell tracking in several ways. The most advantageous characteristics are the superior cellular sensitivity and contrast, and that it allows direct quantification of the SPION content and number of labelled cells at any depth.\textsuperscript{8} These factors contribute to an in vivo cell detection limit down to ~200 labelled cells after implantation,\textsuperscript{26} far surpassing the standard detection limits in MRI.\textsuperscript{260} There has however been evidence to the capability of MRI to track single cells in vivo,\textsuperscript{261-263} but as discussed in section 4.4, with advancements in the sensitivity of MPI scanners and tracers, the theoretical detection limit may also be just a single cell.\textsuperscript{26} Nonetheless, the current properties of MPI permit efficient monitoring of the fate of cell-based therapies and diagnostics, as well as their biodistribution, clearance and retention. There are two primary cell-labelling techniques for cell tracking with MPI.\textsuperscript{264} The first is in situ labelling, where SPIONs are administered intravenously, and phagocytic cells take up the nanoparticles in situ to deliver them to their target site (Fig. 12a). The second is ex vivo labelling, where the relevant cells are collected from the animal, with subsequent incubation/labelling with the nanoparticles in vitro, and injection back into the animal in vivo (Fig. 12b). Currently, many cells and cell-based therapeutics are tracked for biomedical applications, including cancer cells, stem cells for disease treatment, or immune cells for immunotherapeutic cancer treatment.

One of the most important advantages of SPIONs for this application is their biocompatibility. They have a favourable profile of toxicity, and in vitro, generally showing no, or very little short- or long-term effects on cell viability, differentiation or proliferation, in a range of cellular cultures.\textsuperscript{265-268} This indicates the great potential for in vivo applications, especially cell tracking, and the clinical translation of MPI.\textsuperscript{269} One study revealed that rodents can tolerate iron oxide concentrations of up to 3 mmol Fe/kg.\textsuperscript{269} A further Phase I clinical study on the commonly used SPION tracer, ferucarbotran (Resovist), demonstrated safe use of the tracer at doses of 5-40 μmol Fe/kg.\textsuperscript{152} As a further point, SPIONs are completely biodegradable and will remain in the bloodstream until they are metabolised and finally excreted, just like endogenous iron, through the reticuloendothelial system (RES).\textsuperscript{103} Within this system they are primarily cleared by the liver (80%), spleen (5-8%), and the bone marrow (1-2%).\textsuperscript{270}

5.1 Cancer cell tracking. MPI has often been used as a cell tracking modality for immortalised cancer cell lines. Song \textit{et al.}\textsuperscript{271} produced MPI-tailored SPIONs that also possess a good fluorescence imaging performance (Fig. 13a and b).\textsuperscript{12} These nanoparticles enabled efficient ex vivo cell labelling and were sensitive enough to longitudinally track as few as 250 pre-labelled immortal HeLa cells with MPI, following implantation in mice, as displayed in Fig. 13c. Importantly, the nanoparticles did not demonstrate any influence on the cell viabilities, even following a further 48 h of incubation. In another work, Melo \textit{et al.}\textsuperscript{272} compared the quantitative MPI signals and tracking ability of micron-sized iron oxide particles (MPIOs) and ‘gold-standard’ VivoTrax (Magnetic Insight Inc.), for iron-labelled immortal cancer cell lines in a mouse brain.\textsuperscript{28} MPIOs are different from SPIONs typically used for MPI. The particles consist of multiple small cores (~5-10 nm) embedded in a polystyrene matrix.
Resulting from the clustered nature of the cores, they can be regarded as having one large superparamagnetic core (~0.9 μm), which contributes to a very large MPI signal (Fig. 13d). The results show that MPIO-labelled cells can be detected and quantified in the mouse brain model much more readily than VivoTrax-labelled cells (Fig. 13e-g), demonstrating the potential of MPIOs for MPI applications in cell tracking.

Fig. 11 Ferrite nanoparticles, designed for optimal MPI performance. TEM images of different ferrite nanocubes synthesised. The nanoparticles synthesised were (a) Co₀.5Fe₂.5O₄ nanocubes, (b) Zn₀.1Co₀.2Fe₂.7O₄ nanocubes, and (c) Zn₀.2Fe₂.8O₄ nanocubes. (d) PSF demonstrating the MPI signal obtained for each synthesised sample, in comparison to that of references, VivoTrax, and Fe₃O₄ SPIONs. (e) Histograms of the corresponding SNRs for each synthesised sample, the commercial reference, and Fe₃O₄ SPIONs. Reproduced with permission of the Royal Society of Chemistry, from Ref. 13; permission conveyed through Copyright Clearance Center, Inc. TEM images of different SPIONs and Mn-ferrites (MIOs) synthesised. The nanoparticles synthesised were (f) 8 nm SPIONs, (g) 18 nm SPIONs, (h) 8 nm MIOs, and (i) 18 nm MIOs. (j) Hysteresis curves displaying the field-dependent magnetism of each synthesised nanoparticle. (k) Plot of the MPI signal of the SPIONs synthesised in comparison to that of reference, VivoTrax, as a function of sample concentration. Reproduced with permission from Ref. 24. Copyright 2019 American Chemical Society.
Recently, there has been momentum towards studying patient-derived xenograft (PDX) models, supplanting traditional cell lines in cancer research as they better represent the tumor heterogeneity observed in the original tumor.\textsuperscript{271} Studied results are, therefore, more clinically relevant. The ability to produce quality images and longitudinally track and study the fate of PDXs in \textit{vivo} would be very valuable. Knier \textit{et al.} demonstrated the first method for the efficient iron-labelling of PDX cells, which was also the first successful iron-labelling of breast cancer cells derived from patient brain metastases.\textsuperscript{222} Utilising bioluminescence imaging (BLI) to evaluate cell viability, and MPI for detection and quantification of SPION content, they demonstrated sensitive longitudinal tracking of pre-labelled F2-7 PDx cells with MPIOs, where the signals indicate cell viability and tumour proliferation.

In another work developing imaging tools to target and label primary and metastatic lesions, Parkins \textit{et al.} looked to exploit the tumor self-homing phenomenon.\textsuperscript{273} This describes where circulating tumor cells (CTCs) that break away from the primary tumor into the blood stream, preferentially colonise established tumors rather than non-malignant tissues, accelerating metastatic disease. They demonstrate for the first time, the ability of MPI to sensitively detect systemically administered SPION-labelled CTCs and visualise self-homing. This was done in a murine model bearing pre-established human breast cancer lesions, and the labelling demonstrated no detrimental effects on cell viability. The results provide invaluable information about off-target tumor accumulation, as well as the efficiency of CTC infiltration, proliferation, and survival inside tumors, and the potential mechanisms driving self-homing within the body.

5.1.2 Immune cell tracking. A tumour does not just consist of cancerous cells; immune cells (\textit{e.g.}, leukocytes), together with fibroblasts and endothelial cells, form the tumor microenvironment (TME, Fig. 14).\textsuperscript{274} These immune cells interact with tumor cells to influence and modulate the size progression of the tumor. The rapid success and growth of research in cancer immunotherapy has stimulated the need for a method to help determine the location and behavior of immune cells in the TME systematically over time, and to evaluate the efficacy of immunotherapies. Traditionally SPION/H MRI, or \textsuperscript{19}F MRI cell tracking techniques are used.\textsuperscript{275,276} However, these methodologies are considered inadequate. MPI addresses many of the cell tracking needs for immunology and can track SPION-tagged immune cells \textit{in vivo} for the lifetime of the labelled cell.\textsuperscript{26}

The most abundant immune cell type in the TME are tumor-associated macrophages (TAMs). The presence of TAMs has been correlated with many protumoral effects, therefore enhancing and contributing to cancer progression.\textsuperscript{277} Due to their role, they can act as a biomarker to quantifiably monitor both cancer detection and prognosis through imaging. In addition to aiding therapeutic development, the ability to label and image TAMs effectively through SPION phagocytosis will assist in the understanding of their recruitment and infiltration in tumors and could be utilised to predict how aggressive a tumor may be.\textsuperscript{278}
observed (~1 pg Fe/cell). The non-invasive quantitative tracking of adoptively transferred T cell biodistributions will assist in the development of new and effective ACT strategies.

5.1.3 Imaging localised inflammation. A tumour does not just consist of cancerous cells; immune inflammation is a key process in the body’s response to harmful stimuli, including damaged cells and irritants. Hence, the detection and monitoring of localised inflammation could help with the diagnosis of many pathological conditions and the assessment of treatment outcomes following therapeutic intervention. It is characterised by vasodilation, the accumulation of fluid, and the extravasation of immune cells to the inflamed site. Unfortunately, tracking inflammatory processes has traditionally involved low specificity imaging techniques or invasive methods like biopsies.  

Previous MRI studies have already shown the benefits of SPION tracers for inflammation detection and tracking. With MPI, inflammation can be more specifically detected, with a higher sensitivity. Similar to generic cell tracking, there are two primary methods to monitor inflammation with SPIONs. Either the nanoparticles are loaded directly into phagocytic immune cells like macrophages in vivo, with subsequent injection, or they are directly injected, to be engulfed by the cells in situ. Once tagged, these specific cells can be tracked as they migrate and accumulate within regions of localised inflammation.

In a recent example, Mangarova et al. implemented MPI in the ex vivo monitoring of vascular inflammation in abdominal aortic aneurysms (AAAs). AAAs are defined by a weakening and dilatation in the abdominal aorta, usually in the infrarenal portion of the artery, and are currently one of the leading causes of death in developed countries. In this study, the inflammatory response in the aneurysmal wall of an Angiotensin II-infused ApoE −/− mouse model was assessed with in vivo MRI, ex vivo MRI and ex vivo MPS. The ex vivo MRI was performed 24 hours post-intravascular administration of ferucarbotran (Resovist), following 3-4 weeks of Angiotensin II perfusion. The results reveal abundant iron concentration within AAAs following their uptake by macrophages, demonstrating the feasibility of sensitive ex vivo MPI for the detection of vascular inflammation in AAA. These measurements correlated strongly with the results of ex vivo MPS, confirming the inflammatory activity and resulting SPION accumulation in the aneurysmal wall.

The non-invasive tracking of white blood cell (WBC) distributions can also be utilised in the monitoring and diagnosis of inflammation. Traditionally, tracking has been performed using WBCs directly labelled with radionuclides for scintigraphy or SPECT. However, this is not ideal for general screening due to the extensive costs, tracer radioactivity, and high dosage of tracer required. In a very recent work, Chandrasekharan et al. explored for the first time, MPI, for the sensitive and radiation-free tracking of WBCs to sites of inflammation and infection. For this, they used the commercially available antibody-conjugated multi-core Anti-Ly6G SPION (Miltenyi Biotec GmbH, Fig. 15a) for in situ labelling and tracking, which is specific to the Ly6G antigen expressed on neutrophils. Prior to in vivo labelling with these nanoparticles, WBCs were purified from an SPF-grade C57BL/6 mouse and were intravenously injected into C57BL/6 mice. The nanoparticles were conjugated to a multi-core anti-Ly6G antibody with an average size of 45nm, which is significantly smaller than the size of a red blood cell. The Ly6G antigen is expressed on neutrophils and macrophages, making it a suitable target for tracking these cells. The nanoparticles were administered intravenously into C57BL/6 mice and were monitored using MPI and ex vivo MRI.

Fig. 14 Schematic representation of the TME. Reproduced from Ref. 6, with permission from Elsevier.
experiments, the MPI capabilities of this SPION were determined, with Anti-Ly6G demonstrating improved values for MPI spatial resolution (1.26 mm vs 1.49 mm, respectively) and sensitivity (~1.8 times greater) over multi-core VivoTrax (Fig. 15b and c). The in vivo studies were performed in a murine model of lipopolysaccharide-induced myositis, and the nanoparticles were intravenously injected following inflammation induction. MPI was used to image the biodistribution and demonstrated highly sensitive targeting and detection of inflammation at the sites of myositis, with strong contrast-to-noise ratios between 8 and 13 (Fig. 15d).

5.1.4 Stem cell tracking. Stem cell therapies are increasingly recognised as the future of regenerative medicine, demonstrating tissue regeneration potential for a wide variety of conditions. However, for progression of this rapidly growing field to clinical translation, several concerns need to be addressed, most importantly is the poorly understood biological fate and migration of injected or implanted stem cells in the body. Successful therapy requires verification that the stem cells reach and remain at their intended target destination, maintaining viability to ultimately reform functional organs or tissues. It is therefore particularly relevant to monitor and track stem cell biodistribution following administration in vivo, as this would provide valuable information for predicting therapeutic efficacy, evaluating potential risk, and optimising treatment outcomes. Standard preclinical imaging modalities have been used for in vivo stem cell tracking, including MRI and radionuclide imaging. MPI offers significant advantages for longitudinal stem cell tracking over other techniques, which has led to an extensive recent publication history. It has been demonstrated that rat and human adult stem cells uptake SPION tracers, and that they localise within the cytoplasm.

Mesenchymal stem cells (MSCs), which are found in many types of tissue and are multipotent, have shown particularly promising therapeutic results, where MSC-based therapies have shown success in treating many diseases, like stroke, myocardial infarction, and cancer. Importantly, previous MRI studies displayed no decrease in cell viability, proliferation, or differentiation of MSCs after SPION-labelling. However, targeted MSC delivery remains a challenge. Commonly, intravenous deliveries of MSCs become entrapped in lung microvasculature instead of the target tissue. Zheng et al. sought to better understand this observation as they demonstrated, for the first time, the dynamic tracking of MSC administrations in vivo in a rat model with MPI, employing Resovist to label MSCs. The labelled cells were intravenously administered, and the transplantation, dynamic biodistribution, and clearance, were monitored over a 12 day period. An MTT viability assay was carried out, and there was no considerable difference in the cell viabilities of unlabelled and labelled hMSC populations. The MPI images, co-registered with CT, confirm that labelled MSC injections become immediately entrapped in lung tissue and are mostly cleared to the liver within one day. Most significantly though, these results demonstrate that MPI can longitudinally track intravenously administered MSCs safely and quantitatively.

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Fig. 15 The application of anti-Ly6G SPIONs in the imaging of localised inflammation with MPI. (a) TEM image of the anti-Ly6G SPIONs. PSFs demonstrating the MPI signal in saline, and blood, obtained for (b) anti-Ly6G SPIONs, in comparison to that of the reference, (c) VivoTrax. (d) Inflammation MPI images in three different mouse subjects undergoing myositis, with administered anti-Ly6G-SPIONS. Reproduced from Ref. 17 with permission from Ivyspring International Publisher. This is an open access article distributed under the terms of the CC BY License (https://creativecommons.org/licenses/by/4.0/). Copyright 2021, the authors. Article can be found at: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7893534/ No changes were made to the original figure.
In a different work, Nejadnik et al. utilised dual MPI/MRI to track the status of an implanted MSC scaffold over 14 d. The MSCs were labelled with very small FDA-approved ferumoxytol (Feraheme, AMAG Pharmaceuticals) and ferucarbotran (VivoTrax) SPIONs. These labelled MSCs were successfully quantified with MPI at day 1 and 14 (Fig. 16a), with results indicating that MPI was sensitive to changes in cell number of labelled cells at the transplant site over time, with a significant decrease in SPION content observed at the later timepoints (Fig. 16b). Similarly, Sehl et al. implemented a trimodal imaging approach, utilising iron-based \(^1\)H MRI, and quantitative MPI and \(^{19}\)F MRI, to monitor the fate of transplanted ferumoxytol-labelled MSCs and the ensuing inflammation, which is inferred through the tracking of infiltrating macrophages at the transplanted sites in vivo.\(^{19}\) This is the first time these three modalities have been combined to monitor cell populations in vivo. Significantly, the viability of these cells was unchanged following MSC labelling, where a 97% viability was determined pre- and post-labelling. These labelled cells were implanted within the hind limb muscle of C57Bl/6 mice. A perfluorocarbon agent was also administered intravenously for uptake by phagocytic macrophages in situ. The labelled MSCs were detected using \(^1\)H MRI and MPI, and perfluorocarbon-labelled macrophages were detected using \(^{19}\)F MRI. The MPI signal decreased over a 12 day period, which is consistent with the death and clearance of the MSCs, whereas \(^{19}\)F signal persisted (Fig. 16c), suggesting the continuous infiltration of perfluorocarbon-labelled macrophages, and thus inflammation.

In addition to using established SPIONs in the MPI tracking of labelled MSCs,\(^8\) several groups have synthesised their own optimised tracers. Lemaster et al. synthesised hybrid poly(lactide-co-glycolide acid) (PLGA)-based iron oxide nanobubbles (Fig. 17a), labelled with a fluorophore, that show potential as a trimodal imaging tracer (MPI/photoacoustic/US) for MSC labelling.\(^8\) The PLGA coating facilitates the US signal, the iron oxide core enables the MPI signal, and the fluorophore (DiR, 1,1'-dioctadecyl-3,3,3',3'-tetramethylindocarbocyanine iodide) increases the photoacoustic signal (Fig. 17b). They confirmed that there were no adverse effects on cell treatment with nanobubbles in vitro, and hence the nanobubble-labelled cells were injected intramyocardially into live mice for real-time imaging. This multi-modal tracer was shown to track MSCs effectively. In another study, Wang et al. designed a series of specialised cubic tracers with differing edge lengths.\(^{18}\) The tracer with the most beneficial properties for MPI has an edge length of 22 nm, termed CIONs-22 (Fig. 17c). In comparison to the other synthesised tracers of differing sizes, CIONs-22 exhibit good Ms values (Fig. 17d) and high resolution and sensitivity (~2500 cells) for MPI, (Fig. 17e). With their efficient cellular uptake and labelling, these particles enable the long-term and real-time tracking of bone MSCs in vivo for MPI, exhibiting accurate tracking of their migration and distribution pattern once transplanted to hindlimb ischemia mice.

MPI has also demonstrated promise in the monitoring of neural progenitor cell (NPC) and neural stem cell grafts. Progenitor cells are descendants of stem cells that further
differentiate to create specialised cell types. NPCs have been implicated in the treatment of several neurodegenerative disorders, such as brain ischemia, epilepsy, and Parkinson’s and Alzheimer’s disease.\textsuperscript{48,108} The high sensitivity in MPI could facilitate the quantification of dynamic events, such as graft movement, in such brain disease models. In an important \textit{in vivo} study, Zheng et al. implanted SPION-labelled NPCs into the forebrain of rats with forebrain ischemia and monitored the \textit{in vivo} fate of the implanted cells longitudinally with MPI.\textsuperscript{26} First, the NPCs were differentiated from human embryonic stem cells and labelled with ferucarbotran (Resovist). Following injection, it was demonstrated that the labelled cell grafts could be sensitively and quantitatively detected, and that the particles measured nonsignificant signal decay over a period of 87 days (Fig. 18a and b). The authors also demonstrated a detection sensitivity of 200 cells in their custom-built machine. This study therefore exemplifies the feasibility and advantages of long-term sensitive tracking of NPC and stem cell transplants with MPI.

Transplantation of stem cell-derived islet organoids is a promising approach for the treatment of type 1 diabetes.\textsuperscript{301,302} However, there has traditionally been no appropriate imaging technique for the accurate monitoring of graft outcomes after transplantation. Wang et al. demonstrated the use of MPI in monitoring transplanted islets in animal models for the first time.\textsuperscript{303} Pancreatic islets were isolated and labelled with VivoTrax SPIONs before transplantation either under the kidney capsule or in the liver of NOD/scid mice. MPI successfully images and quantifies the islets in these models, post-mortem, at days 1 and 14 following transplantation. At the World Molecular Imaging Congress in 2020, Sun described the feasibility of \textit{in vivo} tracking of transplanted stem cell-derived islet organoids using MPI.\textsuperscript{304} Human induced pluripotent stem cells were differentiated to islet organoids, which were subsequently labelled with VivoTrax following 21 days of differentiation. The labelled organoids were then similarly transplanted into the left kidney capsule of NOD/scid mice, which were examined by MPI/CT up to 28 days post transplantation. There is a strong signal initially detected in the kidney, but the intensity decreases over the study, which is to be expected with organoid clearance. Additionally, they performed an artificial intelligence analysis of the MPI images, to assist with total iron value prediction of the transplanted cells on different days, using an accurate machine learning algorithm established by the same team of researchers.\textsuperscript{305} MPI, assisted by this machine learning algorithm analysis, was shown to accurately monitor islet organoids labelled with SPIONs post transplantation and provide quantitative information on their presence \textit{in vivo}. These results demonstrate the potential for future imaging of cell transplantation and therapy for type 1 diabetes.

5.2 Theranostics

Theranostics, from a clinical and translational viewpoint, refers to an intimate combination of therapeutic and diagnostic interventions. MPI may open unprecedented opportunities for exploring new nanoparticle-based theranostic applications, stemming from the imaging and therapeutic capabilities of SPIONs and the intrinsic properties of MPI itself. In these approaches, diagnostic interventions are most used in combination with targeted MFH or magnetic drug delivery.

5.2.1 Drug delivery. Non-invasive \textit{in vivo} drug release monitoring approaches are highly advantageous. MPI is particularly useful because of the linear quantitation and lack of background signal, providing spatial information which allows the accurate monitoring of therapeutic agent delivery.\textsuperscript{306} This information illustrates the quantity of drug delivered to the target site, as well as the drug dosage distribution within the body. The ability to track drug delivery also allows for real-time adjustment of ensuing doses, ensuring that the dosage at the target site remains within the patient’s therapeutic window (Fig. 19), and that there are subcritical drug concentrations at off-target sites within the patient (e.g., organs).\textsuperscript{15} The
therapeutic window is the dosage range in which the highest therapeutic benefit can be achieved. At suboptimal levels, low non-lethal levels of drug can lead to resistance, where the drug will become less efficient and eventually ineffective (i.e., below the minimum effective dose). At high toxic levels, it can lead to mortality or morbidity (i.e., above the maximum tolerated dose). Drug resistance is the primary reason for the failure of many therapies (e.g., chemotherapy), hence the importance of real-time imaging and adjustments.  

Often, the SPION tracer and therapeutic molecules are co-loaded into the same nanoparticle. In this context, the SPION shell is commonly modified with pharmacochemically active compounds. This can either be done via covalent bonds, or with physical intermolecular interactions, like π-π stacking. SPIONs have been conjugated to many different therapeutics. Most frequently, anti-cancer agents have been employed, from standard chemotherapeutic molecules like doxorubicin (DOX), to more complex therapeutic compounds like siRNA.  

MPi signal intensity increases as they recover their Brownian relaxation (Fig. 20c). This allows accurate quantitation and monitoring of drug release in vitro and in vivo (Fig. 20d and e).  

Fuller et al. also composed magnetic nanocarriers combining high DOX loading and MPI capability (Fig. 20f). Unlike SPNCD, these nanocarriers contain the drug within the hydrophobic core, along with the SPIONs, and are coated with a PEG-block-poly(lactic acid) block copolymer, for water solubility and colloidal stability. The release rate of SPIONS and DOX from the nanocarriers was also dependent on environmental pH (Fig. 20g). A different ‘smart’ agent based on the near-infrared-responsive plasmonic properties of tuned nanostars was developed by Tomitaka et al (Fig. 20h). These engineered nanoparticles are composed of a SPION core and a star-shaped plasmonic shell, made up of high-aspect-ratio gold branches (Fig. 20i). Model drug molecules (TDF, Tenofovir disoproxil fumarate) are bound to the gold shell and can be triggered to release upon near-infrared illumination because of the photothermal effect in the shell (Fig. 20j). Quantitative MPI can be used to monitor the drug release.  

Around 30% of breast cancer tumors will metastasise, most commonly to the liver, lungs, brain and bone. Brain metastases are of a particular concern as they are typically fatal. In the treatment of these brain metastases, targeted therapeutics may not be able to, or will have limited penetration, when attempting to cross the BBB, resulting in reduced treatment efficacy. Extracellular vesicles (EVs) are small particles released by cells and are implicated in numerous biological processes, including metastasis. These particles may cross the BBB and can therefore be exploited to deliver therapeutic agents to brain metastases. However, before further progress in this field, it is important to understand the activity, localisation, and accumulation of EVs during metastatic progression. Toomajian et al. demonstrated the tracking of SPION-labelled EVs within brain metastases using MPI. For in
**5.2.2 Magnetic fluid hyperthermia.** MFH treatment refers to when a suspension of MNPs, administered systemically or locally, is combined with an externally applied AMF to produce heat at a site of nanoparticle accumulation. The heat is generated through the rapid switching of the SPION magnetisation and Brownian directions by the AMF. This hyperthermic effect can lead to tissue damage in the target area surrounding the nanoparticles. To achieve a therapeutic response in tissues using hyperthermia, two treatment approaches are generally employed. In the primary approach, the temperature of the target tissue is increased to ~42 °C and held for a defined time. It is known that healthy tissues have reduced sensitivity to hyperthermia as compared with cancerous cells and may withstand temperatures of 42–45 °C, where cancerous cells undergo apoptosis. This results in denaturation of proteins at the tumour site, and subsequent cell death. In the second approach, the target tissue is completely ablated, causing irreversible damage to the pathological target, at temperatures < 46 °C. Such temperatures lead to the necrosis of cancer cells but may also affect healthy cells, thus hyperthermia-induced apoptosis is preferable. The specific absorption rate (SAR) is used to quantify the effectiveness of a specific SPION for MFH treatment. It is defined as the magnetic power absorbed per unit mass of magnetic material.

The first proposed use of magnetic hyperthermia with iron oxide particles was as early as 1957, yet it remains a prevalent research field today with significant work on the development of novel hyperthermia-tailored nanocarriers. Interest as a therapeutic treatment can be attributed to multiple key advantages, including: no fundamental depth limitation, synergy with many other therapies (e.g., chemotherapies), and internalisation of the ‘fluidic’ heat source. The most frequently used MFH application paradigms are whole-body MFH (Fig. 21a), where homogeneous AMF-generating alternating current (AC) coils scan the whole-body for sites of accumulation, and local MFH (Fig. 21b), where surface AC coils target localised areas of accumulation. Despite the advantages, both approaches face several challenges towards further development and eventual clinical translation. Local MFH struggles with deep-seated treatment and therefore can only be used for treatments at the surface of the body. The most pressing issue for the clinical translation of the more commonly applied whole-body MFH is in the localisation of SPIONs at the target site. Systematically administered SPIONs, even if targeted, will end up in other critical organs like the liver and spleen, as well as the target site. This is challenging since...
accumulation of SPIONs in off-target organs will increase their temperature during AMF application, leading to serious side effects.\textsuperscript{316} Experiments in mice with systemically administered SPIONs and whole-body AMF show elevated liver enzymes, liver and spleen necrosis, and even death.\textsuperscript{322}

Fig. 21 (a) Whole-body MFH targets all tracer, including healthy sites of accumulation. (b) Local MFH targets tracers near the surface only. (c) MPI-MFH can selectively target tracer anywhere, including those deep in the body, while avoiding tracer accumulations in healthy sites. Reproduced with permission of IOP Publishing, from Ref. 23; permission conveyed through Copyright Clearance Center, Inc.

To address these important technical challenges, several research groups have demonstrated theoretically and experimentally the benefits of the combined MFH and MPI approach, MPI-MFH (Fig. 21c).\textsuperscript{178,179,323} The same principles for image generation in MPI, as described in section 2.1, can be modified to spatially select and localise thermal heat deposition to a desired region in biological tissue. MPI uses low sinusoidal excitation frequencies in the drive field, in the order of ~20 kHz, but by exciting SPIONs to higher frequencies of > 300 kHz, heat can be generated.\textsuperscript{321} The particles under the influence of the selection field are locked, and the particles in the FFR are free to rotate, as before, thereby restricting the SPION heating to the FFR alone. Combination with MPI also allows robust treatment planning. It provides precise quantitative imaging of the SPION distribution within the sample before heating, which is essential for quantification of particle accumulation at the target site and thus accurate SAR prediction and appropriate treatment.\textsuperscript{316} As a result of this, and because the MPI-based gradients specifically limit the location of the hyperthermia to only a small, adjustable FFR in the sample, MPI-MFH may overcome the previous limitations with off-target heating experienced with whole-body MPI. The established physics of MPI also benefit MFH by providing high resolution targeting anywhere in the body and without depth limitation, addressing the primary problem encountered with local MFH.\textsuperscript{321} Recently, newer designs for the FFR are being investigated where its size can be varied through alteration of the gradient field strength, thereby allowing correlation of treatment and dosage to the size of the lesion.\textsuperscript{324}

As the physics germane to and exploited by MPI and MFH are similar, the same SPIONs can be used effectively for both. The techniques may also be integrated together in a single device for simultaneous MPI-MFH, that can seamlessly switch between imaging and heating modes through modulation of the AC excitation field magnitude, whilst the subject remains in the scanner. This provides opportunities for real-time diagnostic
study, Bleul et al. demonstrated the use of a micromixer synthesis platform (Fig. 22a) for the size-controlled synthesis of monodisperse single-core SPIONs, with a core diameter of ~30 nm. These particles show promising theranostic capabilities with high signal amplitudes in MPI, and SAR values of up to 1 kW/g(Fe), which exceeds the comparable SAR value of Resovist by more than a factor of three (Fig. 22b). Du et al. also designed monodisperse single-core SPIONs with improved hyperthermia therapy. The hyperthermia performance can also be attributed to SPION surface functionalisation with CREKA, a peptide that actively targets fibrin-fibronectin complexes overexpressed in tumor interstitium, therefore improving delivery uniformity and enabling more effective cancer ablation.

Another factor that may affect the heating performance is the choice of coating material. Jordan et al. first noticed a SAR difference between dextran-coated and aminosilan-coated SPIONs, with aminosilan particles having a 1.2-fold greater SAR in identical particles. Liu et al. then performed a systemic analysis of the SAR produced by SPIONs coated in different PEG molecular weights, observing a SAR increase with thinner coatings. This is attributed to the dominance of Brownian relaxation based heat losses. However, it is noted that reducing the thickness may also affect the colloidal stability of the SPIONs, which detrimentally affects SAR. Differently, shape anisotropy in SPIONs has also been shown to improve hyperthermia performance, as is described in section 4.1.

Khurshid et al. reported a 1.4-fold improvement in the particle SAR for cubic SPIONs over spherical SPIONs. Bauer et al. also reported the high performance of cubic SPIONs, this time selectively doped with anisotropic zinc, increasing the overall anisotropy. They observed a 2-fold enhancement of MPI signal and a 5-fold improvement in SAR, in comparison with equivalent undoped single-core spherical SPIONs.

5.2.3 Combinatorial therapy. The theranostic approaches described above can also be extended for combinatorial therapy, in which the actuated release of drugs from a nanocarrier can be achieved in combination with hyperthermia, providing great spatial and temporal control of release. The amount of drug release can be tuned via alteration in the AMF field strength, and negligible release takes place when the particles are not subjected to an AMF. Maruyama et al. developed magnetic nanocarriers that encapsulate SPIONs and DOX, based on thermosensitive liposomes. The results indicated the usefulness of MPI for both in vitro and in vivo studies for monitoring drug release through SPION release, following disruption of liposomal membranes by AMF-induced hyperthermia. However, these authors characterised the MPI performance of their liposomes using a prototype MPI scanner (400 Hz, 16 mT) that operates at higher drive field amplitudes and much lower drive field frequencies than most commercial scanners, and this influences the MPI resolution and signal intensity. Rost et al. demonstrated that the MPI performance of similar SPION-encapsulating liposomes is maintained using the commercially available MOMENTUM scanner (Magnetic Insight Inc., scan at 45 kHz, 16 mT). In a different work, Fuller et al. developed a drug co-polymer conjugate encapsulating semispherical 18 nm single-core SPIONs and provided the proof-of-concept for spatially controlled drug release using an FFR, in this case following the breakage of a thermally-labile Diels-Alder bond. The release can potentially be monitored with MPI and activated numerous times, and the magnetic field gradients can be used to spatially control the region of release, thus preventing damage to off-target tissue.

5.3 Perfusion imaging

Perfusion imaging is a technique whereby the passage of a tracer is monitored continuously through the capillary tissue bed, capturing any temporal changes in the tracer. It has seen extensive application in the diagnosis and detection of various pathophysiology’s related to blood perfusion and vascular changes. Steaming from the short scanning time and high temporal resolution of MPI, it has become possible to monitor perfusion and blood-flow in real-time. The speed at which images are captured would substantially benefit diseases for which a rapid assessment of the vasculature and perfusion are mandatory for the treatment. Another beneficial property of

Fig. 22 MPI-tailored single-core SPIONs, designed for MPI-MFH. (a) Photograph of the micromixer system used for the synthesis of 30 nm monodisperse single-core SPIONs. (b) SAR values of the synthesised SPIONs as a function of frequency (f), in comparison to that of the reference, Resovist. Reproduced with permission of the Royal Society of Chemistry, from Ref. 9; permission conveyed through Copyright Clearance Center, Inc.
MPI for perfusion imaging is the high contrast-to-noise, permitting higher accuracy imaging that compares favourably to the traditionally applied techniques of contrast-enhanced CT or MRI, which produce unreliable perfusion images.335

Neuropathological diseases such as ischemic stroke, and traumatic brain injury (TBI) are severe conditions requiring immediate medical attention and extensive monitoring following treatment. Specifically stroke, with 17 million cases worldwide every year, is one of the leading annual causes of death and disability.171 Rapid high quality cerebral perfusion imaging is fundamental in successful stroke diagnosis and management, as it allows for early assessment of penumbra volume and location, as well as in predicting which patients may benefit from cerebral revascularisation therapies.236 The low temporal resolution of perfusion MRI and CT can cause variations in the penumbra volume estimate, which may affect optimal stroke treatment.337,338 In a pivotal study, Ludewig et al. demonstrated the potential of MPI in the real-time perfusion imaging and diagnosis of acute stroke in a rodent model.171 Cerebral ischemia was induced in the internal carotid artery of C57BL/6 mice, and within seconds and a single injection of long-circulating LS-008 SPIONs, cerebral perfusions could be assessed by MPI. The signal lowered in the ischemic hemisphere, allowing precise detection of an ischemic stroke within a few cubic millimetres. The SPIONs persist in the bloodstream for hours following injection, without any leakage into the brain interstitium.

Long-circulating tracers are crucial in most perfusion imaging applications, enabling blood flow and blood volume studies. Orendorff et al. exhibited their importance in the imaging of TBI, where a closed-head rat model was monitored longitudinally to study cerebral bleeding caused by an impact, as well as changes in the blood pool as the wound heals.170 This work utilised single-core LS-13 nanoparticles (LodeSpin Labs) that show a circulation half-life of approximately 4-6 hours. Such times allowed infiltration into the blood pools in the interstitial space, and thus successful monitoring of TBI with MPI over long time scales, without significant signal loss.

In addition to TBI imaging, MPI has shown great promise for various other blood pool imaging applications, including the detection of gastrointestinal (GI) bleeding. GI bleeding is a serious clinical issue associated with the haemorrhaging of organs in the digestive system, yet its diagnosis remains very challenging. Traditionally, diagnosis is achieved using scintigraphy techniques in which a RBC is tagged using a radioisotope like 99mTc.339 However, the bleeding site must be located quickly for rapid diagnosis and intervention, and the long hot chemistry preparation times for 99mTc RBCs are not ideal.340 An accurate and timely diagnosis method is therefore essential to reduce morbidity and mortality, and for the determination of transfusion requirements. Yu et al. demonstrated that MPI could be utilised for the safe and quantitative detection of GI bleeding, without any hot chemistry preparation time.58 This work was applied to a murine model that is genetically predisposed to polyph development in the GI lumen (Age<sup>Min</sup>/), with subsequent heparin injection to induce acute GI bleeding. Following the injection of long-circulating MPI-tailored SPIONs (single-core LS-017, LodeSpin Laboratories) through the tail vein, dynamic MPI projection images could be analysed to show tracer accumulation in the lower GI tract. These images illustrate that acute GI bleeding could be detected with high sensitivity using MPI in vivo, with bleed rates as slow as 1 to 5 μL/min.

Accurate imaging of lung perfusion is required for the diagnosis of conditions like pulmonary embolism (PE), as many patients experience mild or non-specific symptoms. PE carries a 30% mortality rate when left untreated, however following time-sensitive detection and treatment, this mortality rate may reduce to ~8%.27 Zhou et al. reported the first MPI-based lung perfusion study in vivo in a healthy rat model.27 In this work, they utilised novel multi-core SPIONs that are conjugated to macroaggregated albumin (MAA), aptly termed MAA-SPIONs (Fig. 23a). These are administered intravenously where the particles effectively pass through the lung capillaries after 15 min as the MAA-SPION size is larger than that of the capillary diameter, becoming entrapped in the lung capillary bed (Fig. 23b). This enables the accurate and reliable diagnosis of any vascular related pulmonary defects. In comparison, non-conjugated SPIONs alone cannot target the lung and are instead immediately cleared to the liver and spleen (Fig. 23c). 3D MPI lung perfusion images are comparable in quality to that of traditionally used scintigraphy and SPECT with MMA-conjugated 99mTc, whilst also demonstrating advantages over many modalities due to no air-tissue interface artifacts nor ionising radiation. In a different study, Tay et al. showed that inhaled multi-core SPIONs can be tracked and quantified with MPI in mice, with accuracies comparable to radiolabelled aerosols.159 When taken with the results from Zhou et al.,27 the authors demonstrated a proof-of-concept for MPI-based perfusion-ventilation mapping.51 This mapping has been applied clinically using other modalities, for the diagnosis of PE and in preoperative evaluation of the lungs.

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5.3.1 Cancer detection and screening. The rate of cancer curing almost directly correlates with the stage at which it is diagnosed, meaning that if it was diagnosed and provided with appropriate treatment at an earlier stage, the cure rate would increase significantly.\textsuperscript{342} Biomedical imaging plays a key role in all phases of clinical cancer management. The most important challenge in cancer imaging is the ability to reliably distinguish the tumor from healthy tissue. Current anatomical imaging techniques, like MRI and CT, are useful in detecting changes in the tissue architecture that generally accompany cancer, but the tumor contrast generated is not sufficiently different from healthy tissues for confident diagnoses.\textsuperscript{343} This is most true for metastatic and diffuse tumors. MPI displays excellent contrast, and thus shows great promise for earlier cancer detection and staging than other current imaging methods.

For tumour visualisation, SPIONs must first become localised within the tumor. Currently this process has relied on the enhanced permeability and retention (EPR) effect, otherwise known as ‘passive targeting’.\textsuperscript{344,345} This is a phenomenon observed in cancer where the improper alignment of endothelial cells, poor drainage of the lymphatic system, and leaky vasculature of a growing tumor allow larger sized therapeutic agents, like SPIONs or antibodies, access to the interstitial portion of the tumor, to some extent (Fig. 24). This leads to substantial non-targeted accumulation of such agents, at concentrations several folds higher than in the plasma.\textsuperscript{344}

Yu et al. demonstrated the EPR phenomenon with MPI for the detection of triple negative breast cancer xenograft tumors in an immune deficient murine model.\textsuperscript{3} Long-circulating LS-008 tracers were administered intravenously through the tail vein, and MPI exhibited quantitative visualisation of the tracer dynamics in the tumor over time, showing initial EPR wash-in into the tumors, and delayed wash-out 48 hours later. Notably, the tumor can clearly be detected with MPI, even with large cardiac and liver signals, highlighted with a tumor-to-background ratio of up to 50.

There have also been various examples of non-targeted SPIONs that have been specifically developed to exploit the EPR effect for MPI, demonstrating significant increases in accumulation and MPI signal in tumors based on passive targeting alone.\textsuperscript{10,28} Additionally, to improve the specificity and uptake of SPIONs for cancer detection and imaging, and as active targeting is not possible in all tumors, there have also been many cases of researchers binding tumor biomarker targeting moieties to SPIONs for MPI.\textsuperscript{4,24,30} This active targeting should further improve the capability of MPI for early cancer diagnosis.

5.3.2 Angiography. The rate of cancer curing almost directly correlates with the stage at which it is diagnosed, meaning that if it was diagnosed and provided with appropriate treatment at an earlier stage, the cure rate would increase significantly.\textsuperscript{342} Biomedical imaging plays a key role in all phases of clinical cancer management. The most important challenge in cancer imaging is the ability to reliably distinguish the tumor from healthy tissue. Current anatomical imaging techniques, like MRI and CT, are useful in detecting changes in the tissue architecture that generally accompany cancer, but the tumor contrast generated is not sufficiently different from healthy tissues for confident diagnoses.\textsuperscript{343} This is most true for metastatic and diffuse tumors. MPI displays excellent contrast, and thus shows great promise for earlier cancer detection and staging than other current imaging methods. One of the most compelling applications of MPI is in safe angiography. MPI angiograms allow visualisation of the vasculature. This is of great importance in the diagnostics of many vascular diseases, potentially detecting abnormalities such as deep vein thrombosis, aneurysms, or stenosis. The earliest in vivo experiment carried out using MPI was the real-time 3D imaging of the blood-flow in a beating mouse heart.\textsuperscript{50} These images displayed large vessels like the vena cava, as well as the cardiac chambers. However, the heart and vena cava of mice have been scanned numerous times since then, with more sophisticated MPI-tailored tracers,\textsuperscript{219} and with various improved MPI systems, incorporating advances in MPI technology.\textsuperscript{219,234,346} With these advances, there is an obvious improvement in angiographic image quality.

Kaul et al. demonstrated the improvement in angiographic imaging quality of single-core LS-008, a tracer tailored towards angiography and blood pool imaging in MPI, over MRI-tailored multi-core Resovist.\textsuperscript{3} For in vivo experiments, the tracers were injected into equivalent healthy FVB mice. Whilst both tracers could visualise the propagation of the bolus through the inferior vena cava (Fig. 25a and b), LS-008 clearly displayed fewer temporally fluctuating artifacts, and the signal modulation in the caval vein, resulting from periodic cardiac and respiratory motion, could be obviously depicted. Additionally, the aorta was clearly distinguishable from the caval vein (Fig. 25c), and several further vessel structures and processes could be observed and monitored that were missing with Resovist. These results demonstrate that regarding the quality of delineation and number of visualised vessels, tailored LS-008 outperforms Resovist in angiographic and perfusion MPI.

In a similar study, Mohtashamolpatshahi et al. compared the in vivo imaging quality of angiographic MPI for newly developed multi-core particles (MCP 3),\textsuperscript{207} and Resovist, in the inferior vena cava and aorta.\textsuperscript{347} The tracers were administered intravenously into the tail veins of rat models at doses of 0.1,
This study was performed in the inferior vena cava of healthy FVB mice, and the images obtained with MPI were compared with those from MRI. Analysis revealed good agreement between the MPI and MRI images. The MCP 3 MPI images demonstrated a significantly higher image quality than those with Resovist. Following administration of MCP 3 at dosages of 0.1, and clinically acceptable 0.05 mmol Fe/kg, morphological features such as vessel lumen diameters of the inferior vena cava and abdominal aorta could be assessed, which was not possible with Resovist images at any dosage. Additionally, there are fewer severe background noise artifacts than with Resovist images. It can therefore be said that MCP 3 increased the visibility of vessel lumens in vivo in MPI, towards the possible detection of vascular abnormalities.

Differently, Kaul et al. demonstrated the potential of MPI for quantifiable measurements of in vivo blood-flow velocities. This study was performed in the inferior vena cava of healthy FVB mice, and the images obtained with MPI were compared with those from MRI. Analysis revealed good agreement between the in vivo velocities, with MRI at 4.0 ± 1.5 cm/s and MPI at 4.8 ± 1.1 cm/s. They also performed an in vitro study, where a phantom setup mimicking the flow within the inferior vena cava concluded that velocities of up to 21 cm/s can be captured by MPI as they occur.

Recently, Molwitz et al. obtained the first angiographic MPI images from organs of human-size. This work was performed using a multi-modal ex vivo porcine kidney perfusion system compatible with MPI and well-established magnetic resonance angiography (MRA). This developed system practically evaluates MPI’s potential for human-sized angiography, which is especially relevant as current angiography-suited MPI scanners still suffer from subject size and spatial resolution restrictions. All the visible vessels in MRA and MPI were compared, and where 33% of all vessels imaged in MRA were visible in MPI. This difference is likely a result of the restricted spatial resolution in MPI. Despite these current limitations, this work does demonstrate MPI’s capability for detecting vessels within a single human-sized organ and will be useful for improving MPI’s angiographic potential.

### 5.3.3 Nanowarming of cryopreserved organs

The shortage of available donated organs is one of the greatest crises facing modern medicine today, with many patients dying whilst waiting for potentially lifesaving organs. One of the reasons donated organs never reach a recipient is that they exceed their preservation window following removal. There is therefore a dire need for better-quality organ preservation strategies that enable the biobanking of organs and extend their preservation times. Through this, compatible donor-to-recipient matching would be possible, allowing extensive treatment planning and decreased recipient organ rejection. One method of particular interest for storage is cryopreservation, where the organ is rapidly cooled to extreme low temperatures. However, this technique is not without its challenges. Cryopreservation agents are toxic and require rapid cooling to vitrify and even more rapid warming to avoid devitrification. Currently applied warming modalities lead to temperature gradients and nonuniform rewarming, as well as thermal stresses that can damage larger organs. These warming limitations can be overcome through a nanowarming method, which enables fast and uniform volumetric heating of cryopreserved tissues. In nanowarming, SPION-containing magnetic cryopreservation agents (mCPAs) are uniformly perfused within the target tissue, prior to vitrification. The SPIONs can then be heated when the tissue is required through application of an external AMF, which is capable of penetrating the entire tissue volume without attenuation.

Chiu-lam et al. reported the first successful example of the cryopreservation and nanowarming process in a whole organ, using an mCPA solution. 3D MPI was chosen as an appropriate non-invasive and quantitative technology to evaluate the distribution of the perfused mCPA within the organ. MPI demonstrated that a stable mCPA solution containing a VSSS cryopreservation agent, and especially formulated non-toxic SPIONs that have exceptional stability within VSSS, can uniformly perfuse whole rat hearts. Once this uniformity was determined, the hearts were successfully vitrified, cryopreserved in liquid nitrogen for up to 1 week, and rapidly nanowarmed in a uniform fashion to room temperature using an AMF. These formulated SPIONs demonstrated ultrafast nanowarming rates (> 320 °C/min), that can be controlled with AMF amplitude. To complete the process, the SPIONs were subsequently perfused out, at a 95% success rate as determined by MPI. In this work, there was no evidence of macroscopic damage or stress to the hearts after the processes of vitrification, cryopreservation, and nanowarming. These results support the potential of nanowarming, utilising MPI to visualise mCPA distribution, as a strategy for biobanking tissues for transplantation, potentially significantly enhancing the availability of viable donor organs.
6. Multi-colour MPI

Recently, it has been demonstrated that MPI can take advantage of SPIONs with differential magnetic relaxation or harmonic response behaviors to generate multiple contrasts, enabling ‘multi-colour’ imaging and the ability to measure distinct signals corresponding to a specific particle subtype.\(^{353,354}\) This enablement of MPI to simultaneously measure and separate the signal generated by different SPION types would be useful for many applications in the fields of intervention, therapy, and medical imaging.

Multi-colour imaging was first experimentally realised using x-space and SFR-based approaches. Hensley et al. demonstrated that x-space reconstruction is capable of differentiating between the relaxation behaviors of different SPIONs, with multiple measurements at different drive field amplitudes.\(^{355}\) Rahmer et al. demonstrated that different SPIONs could be differentiated based on the differences in their harmonic response through the SFR approach, where an extensive calibration procedure was performed separately for each SPION type.\(^{356}\) Through the x-space reconstruction process, the work was first extended to in vivo imaging, where the images successfully differentiate between MAA-SPION tracers in the lungs and Chemicell tracers in the liver of a murine model.\(^{356}\) However, the x-space and SFR approaches rely on extensive calibrations or measurements to capture the differences in the relaxation behaviors or harmonic responses of the nanoparticles. To negate this, Muslu et al. proposed a calibration-free method for x-space MPI, where no prior information about the SPIONs is required.\(^{357}\) This technique can generate a multi-colour relaxation time constant map of different SPION types from a single back and forth scan of the FOV, at a single drive field amplitude.

There has been significant progress in using multi-colour MPI for catheter tracking during vascular interventions.\(^{358}\) In such applications, one SPION type is injected into the blood stream for vessel visualisation, whilst a different type of SPION is used to coat a catheter. Multi-colour MPI can discriminate between the particles impregnated in the catheter and those flowing in the vessels. In a recent work, following the staining of a blood vessel phantom and catheter with different SPIONS, Rahmer et al. demonstrated how 3D real-time multi-colour MPI feedback with an online reconstruction can be used to steer the catheter into a desired vessel phantom, following implementation of an external magnetic field.\(^{359}\)

With multi-colour MPI, it is not only possible to distinguish between particles with different characteristics, but also identical particles in different environments.\(^{360}\) Multi-colour MPI can therefore be used to map information relating to the local microenvironment. There are many potential environmental factors where changes could produce different MPI signals. Essentially any factor that causes change in the particle spectrum could form a basis for discrimination. For instance, the temperature mapping capability of multi-colour MPI is well documented, with identical SPIONS in differently heated environments able to be differentiated.\(^{361,362}\) Recently, Paysen et al. were even able to discriminate between free and cell-bound SPIONs by their relaxation behavior, forming multi-colour images (Fig. 26a and b).\(^{2}\) They quantified the dynamic changes that occur when free SPIONs come into contact with and are internalised by cells in vitro over time, without any damage to the cells.

There has been sustained work in viscosity mapping with multi-colour MPI.\(^{363}\) Certain diseases, like cancer and atherosclerosis, are known to significantly increase the levels of cellular viscosity. These diseases can be potentially probed with MPI, through tissue viscosity measurements at the locations of SPION accumulation.\(^{22}\) Möddel et al. reported a method that allowed them to experimentally determine the viscosity of a small sample using a novel multi-colour reconstruction approach.\(^{364}\) This approach was adapted from a previously reported method for temperature mapping with MPI.\(^{362}\) A series of samples with differing viscosities were prepared with glycerol/distilled water mixtures of varying proportions and a fixed concentration of Resovist. Using these samples, they were able to determine the viscosity of the particle environment within a range of 1.0-51.8 mPa s, with a methodological error of 6%. More recently, Utkur et al. demonstrated the results of relaxation-based multi-colour MPI for viscosity mapping.\(^{22}\) An imaging phantom that contained SPION samples at five different viscosity levels was prepared (Fig. 27a), again using water/glycerol mixtures of varying proportions. The resulting samples covered the biologically relevant viscosity levels, with values between 0.89 mPa s and 5.04 mPa s. Using the relaxation time constant estimation technique outlined by the same group,\(^{367}\) they showed that relaxation-based experiments can distinguish multi-core SPIONs within the biologically relevant viscosity range (Fig. 27b-d).

7. Clinical translation

Commercial pre-clinical MPI scanners are only available through Magnetic Insight Inc. and Bruker GmbH. Pre-clinical system
specifications are slowly improving as researchers have been highly active in designing improved software for image acquisition and reconstruction.35,37,39,40,365-373 For example, Gladiss et al. recently demonstrated the advantages of a hybrid system matrix for efficient calibration and image reconstruction.374 To scale-up MPI hardware to human-size, there are several notable safety considerations and ergonomic challenges that must be overcome. Most notably, issues regarding potential allergic reactions to SPION injections,375-377 and the higher cost and power consumption requirements for the larger field amplitudes needed for sufficient image resolution at a clinical scale.80 Initially, tissue heating and peripheral nerve stimulation resulting from time-varying magnetic fields was also thought be an issue. However, studies have demonstrated that the physical properties of the magnetic fields required for scanning human-sized volumes in MPI can be tuned as to not cross the limit at which these effects are generated, without having a negative effect on performance.378-380

Despite these obstacles, there has been sustained work on the translation and hardware scale-up.32,43,79,235,381,382 In the design of a functional MPI brain imager, Mason et al. developed simulation studies that demonstrated promising capabilities for human-scale systems.383 In an alternative approach Graeser et al. successfully presented a human-sized MPI hardware set-up, tailored for brain applications, that has low technical requirements for fast and flexible operation in a clinical environment.384 For a different application, Mason et al. also developed a highly sensitive small-bore 2D FFL MPI projection imager, that can rapidly image the distribution of tumors in excised breast tissue, from intravenously injected SPIONS, during breast-conserving surgery treatments for breast cancer.385 This work shows the potential for MPI as a clinical solution to issues with positive margins often seen in such surgeries. Another vital area for focus in advancement towards clinical MPI is in the development and commercialisation of SPIONs tailored towards MPI physics. Currently, no such SPIONs have been approved for clinical treatment, yet it has already been demonstrated that enhanced MPI performance can be realised through specific SPION design, with improvements in, for example, MPI spatial resolution and sensitivity.386-388 Notably, further optimisation of the SPIONs for spatial resolution can be traded-off for lower MPI gradients, lowering the overall cost of MPI implementation and easing the scale-up of hardware to human size.230 With different cores, coatings, and the potential conjugation of targeting and therapeutic moieties, it is anticipated that SPIONs will be designed for many different applications and disease processes. Particularly, there has been increased interest in the design of smart multifunctional nanoparticle assemblies for drug delivery applications.15,21,25,181,333,334 This work could be extended to designing activatable nanomaterials where the breakdown and release of SPIONs is triggered by enzyme activation. Such particles could be used to measure enzymatic levels in an in vivo area through enzymatic sensing.

8. Conclusions and perspectives

MPI is an emerging radiation-free imaging modality that utilises sensitive, safe, and biocompatible SPIONs as its tracing material. It demonstrates totally unambiguous depth-independent detection of these tracers, negligible background signal, and superb image sensitivity, which enables an essentially infinite contrast and the production of highly specific images with very low detection limits. MPI is also truly linearly quantitative, demonstrating an almost perfectly linear relationship between MPI signal and iron content, allowing estimation of SPION concentration in target tissues, based on just the signal intensity. This quantitation can even be demonstrated in regions that are challenging for other modalities, like the lungs or bone marrow. These great advantages enable a variety of applications of clinical relevance, as well as unprecedented new applications that were inaccessible using other modalities. Recently, impressive progress has been made in pre-clinical animal studies, in areas such as MFD, drug delivery, inflammation imaging, cell tracking, and perfusion imaging.

For advancing the biomedical applications of MPI, the design of optimal SPIONs and rigorously controlled syntheses have also become important topics of research. Great progress has been made, with many new SPIONs, and other MNPs, demonstrating impressive improvements in general MPI performance,5,10,14 and performance in specific MPI applications.15,191,219 Critical parameters to control include the size and shape of the core, the specific functionalisation, and of utmost importance, assessing whether a single- or multi-core is appropriate for an intended application.

Despite the sustained research, there is plenty of space for the further expansion of MPI. The development of multi-colour MPI and its reconstruction approaches would drive the modality forward, extracting information relating to the local nanovironment that could be used in therapy and diagnostics. Viscosity mapping for example, should be applied in the detection of blood coagulation or diseases that are related to viscosity changes. In addition, MPI has been only
recently applied to the nanowarming of cryopreserved organs, where it is implemented to quantitatively assess SPION loading in an organ before vitrification and after nanowarming. To further this, nanowarming could potentially be coupled with the capability of MI to control the location of SPION heating. This would facilitate great control over the resulting temperature distribution during the rewarming process. There has also been encouraging progress in the design and development of hybrid scanning systems where MI is merged with complementary imaging techniques to improve imaging quality, such as in MI/MRI and MI/CT.

With multiple promising approaches for clinical translation, improved SPION design, and sustained research into the great number of biomedical applications of clinical promise, it can be anticipated that MI will be a crucial complimentary clinical diagnostic and therapeutic tool in the near future.

Author contributions
Stanley Harvell-Smith: Conceptualization, Investigation, Visualization, Writing – original draft, Writing – review & editing. Le Duc Tung: Writing – review & editing. Nguyen T. K. Thanh: Conceptualization, Supervision, Writing – review & editing.

Conflicts of interest
There are no conflicts to declare.

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References
1. L. Glog, M. Mehdipour, M. Ulanova, K. Mariandry, M. A. Nichol, D. J. Hernandez-Castillo, J. Gaudet, R. R. Qiao, J. Zhang, M. Nelson, B. Thierry, M. A. Alvarez-Lemus, T. T. Tan, J. J. Gooding, N. Braidy, P. S. Sachdev and R. D. Tilley, Chem Commun., 2020, 56, 3504-3507.
2. H. Paysen, N. Loewa, A. Stach, J. Wells, O. Kosch, S. Twamley, M. R. Makowski, T. Schaeffter, A. Ludwig and F. Wiekhorst, Sci. Rep., 2020, 10, 1922.
3. M. G. Kaul, T. Mumert, C. Jung, J. Salamon, A. P. Khandhar, R. M. Ferguson, S. J. Kemp, H. Ittrich, K. M. Krishnan, G. Adam and T. Knopp, Phys. Med. Biol., 2017, 62, 3454-3469.
4. H. Arami, E. Teeman, A. Troksa, H. Bradshaw, K. Saatchi, A. Tomitaka, S. S. Gambhir, O. U. Hafeli, D. Liggitt and K. M. Krishnan, Nanoscale, 2017, 9, 18723-18730.
5. S. K. Avugadda, S. Wickramasinghe, D. Niculaes, M. Ju, A. Lak, N. Silvestri, S. Nitti, I. Roy, A. C. S. Samia and T. Pellegrino, Nanomaterials (Basel), 2020, 11.
6. P. Prajapati and D. W. Lambert, J. Bone Oncol., 2016, 5, 128-131.
7. Y. Dai, C. Xu, X. Sun and X. Chen, Chem. Soc. Rev., 2017, 46, 3830-3852.
8. J. E. Lemaster, F. Chen, T. Kim, A. Hariri and J. V. Jokster, ACS Appl. Nano Mater., 2018, 1, 1321-1331.
9. R. Bleul, A. Baki, C. Freeese, H. Paysen, O. Kosch and F. Wiekhorst, Nanoscale Adv., 2020, 2, 4510-4521.
10. G. Song, M. Kenney, Y. S. Chen, X. Zheng, Y. Deng, Z. Chen, S. X. Wang, S. S. Gambhir, H. Dai and J. Rao, Nat. Biomed. Eng., 2020, 4, 325-334.
11. N. H. Paysen, A. Lahiji, K. Yerneni and H. E. Daldrup-Link, Mol. Imaging Biol., 2019, 21, 465-472.
12. G. Song, M. Chen, Y. Zhang, L. Cui, H. Qu, X. Zheng, M. Wintermark, Z. Liu and J. Rao, Nano Lett., 2018, 18, 182-189.
13. N. Silvestri, H. Gavil, P. Guardia, R. Brescia, S. Fernandes, A. C. S. Samia, F. J. Teran and T. Pellegrino, Nanoscale, 2021, Advance Article.
14. Z. W. Tay, S. Saviwala, D. W. Hensley, C. Colson, B. D. Fellows, X. Zhou, Q. Huynh, L. Yu, L. Zhong, P. Chandrasekharan, S. M. Rivera-Jimenez, C. M. Rinaldi-Ramos and S. M. Conolly, Small Methods, 2021, 2100796.
15. X. Zhu, J. Li, P. Peng, N. H. Nassab and B. R. Smith, Nano Lett., 2019, 19, 6725-6733.
16. M. Unni, A. M. Uhl, S. Saviwala, B. H. Savitzky, R. Dhavalikar, N. Garraud, D. P. Arnold, L. F. Kourkoutis, J. S. Andrew and C. Rinaldi, ACS Nano, 2017, 11, 2284-2303.
17. P. Chandrasekharan, K. L. B. Fung, Y. X. Zhou, W. Cui, F. Colson, D. Mai, K. Jeffris, Q. Huynh, C. Saayuy, L. Kabuli, B. Fellows, Y. Lu, E. Yu, Z. W. Tay, B. Zhong, L. Fong and S. M. Conolly, Nanotheranostics, 2021, 5, 240-255.
18. Q. Y. Wang, X. B. Ma, H. W. Liao, Z. Y. Liang, F. Y. Li, J. Tian and D. S. Ling, ACS Nano, 2020, 14, 2053-2062.
19. O. C. Sehl, A. V. Makela, A. M. Hamilton and P. J. Foster, Tomography, 2019, 5, 367-376.
20. S. Liu, A. Chiu-Lam, A. Rivera-Rodriguez, R. DeGropp, S. Saviwala, N. Sarna and C. M. Rinaldi-Ramos, Nanotheranostics, 2021, 5, 348-361.
21. E. G. Fuller, G. M. Scheutz, A. Jimenez, P. Lewis, S. Saviwala, S. Liu, B. S. Sumerlin and C. Rinaldi, Int. J. Pharm., 2019, 572, 118796.
22. M. Utkur, Y. Muslu and E. U. Saritas, Phys. Med. Biol., 2017, 62, 3422-3439.
23. D. Hensley, Z. W. Tay, R. Dhavalikar, B. Zhong, P. Goodwill, C. Rinaldi and S. Conolly, Phys. Med. Biol., 2017, 62, 3483-3500.
24. Y. Du, X. Liu, Q. Liang, X. J. Liang and J. Tian, Nano Lett., 2019, 19, 3618-3626.
25. A. Tomitaka, H. Arami, A. Ahmadianvand, N. Pala, A. J. McGoron, Y. Takemura, M. Febo and M. Nair, Sci. Rep., 2020, 10, 11015.
26. B. Zheng, T. Vazin, P. W. Goodwill, A. Conway, A. Verma, E. U. Saritas, D. Schaffer and S. M. Conolly, Sci. Rep., 2015, 5, 14055.
27. K. K. Melo, A. V. Makela, A. M. Hamilton and P. J. Foster, bioRxiv, 2020, preprint, DOI: 10.1101/2020.07.12.197780.
28. A. Tomitaka, O. K., K. Nishimoto, H. Arami, Y. Takemura and M. Nair, Nanoscale, 2019, 11, 6489-6496.
29. B. Gleich and R. Weizenecker, Nature, 2005, 435, 1214-1217.
243 Z. Zhao and C. Rinaldi, presented in part of World Molecular Imaging Congress (WMIC), Virtual, October 2020.
242 C. Saayujya, presented in part of World Molecular Imaging Congress (WMIC), Montreal, September 2019.
241 C. Colson, Z. W. Tay, K. L. B. Fung, D. W. Hensley, S. Savliwala, S. Horvat, P. Vogel, D. L. White, P. Jacobs and J. Lewis, Compton, D. L. White, P. Jacobs and J. Lewis, Phys. Med. Biol., 2016, 60, 145101.
240 Z. W. Tay, PhD Thesis, UC Berkeley, 2018.
239 W. Paysen, O. Kosch, J. Wells, N. Loewa and F. Wiekhorst, Phys Med Biol, 2020, DOI: 10.1088/1361-6560/abc364.
238 P. W. Goodwill, E. U. Saritas, L. R. Croft, T. N. Kim, K. M. Krishnan, D. V. Schaffer and S. M. Conolly, presented in part of World Molecular Imaging Congress (WMIC), Virtual, October 2020.
237 C. Shasha, E. Teeman and K. M. Krishnan, R. Dhavalikar and C. Rinaldi, presented in part of World Molecular Imaging Congress (WMIC), Virtual, October 2020.
236 P. W. Goodwill, E. U. Saritas, L. R. Croft, T. N. Kim, K. M. Krishnan, D. V. Schaffer and S. M. Conolly, presented in part of World Molecular Imaging Congress (WMIC), Montreal, September 2019.
235 H. Paysen, O. Kosch, J. Wells, N. Loewa and F. Wiekhorst, Phys Med Biol, 2020, DOI: 10.1088/1361-6560/abc364.
234 M. Graeser, T. Knopp, P. Szwargulski, T. Yoneda, I. C. MacDonald, A. F. Chambers, B. K. Rutt and P. J. Foster, Magnetic Resonance in Medicine, 2021, 86, 130-137.
233 R. Dhavalikar and C. Rinaldi, Journal of Applied Physics, 2014, 115.
232 K. Rezwan and R. Dringen, Acta Biomater., 2014, 10, 11548-11551.
231 J. Yu, C. Yang, J. D. S. Li, Y. C. Ding, L. Zhang, M. Z. Yousaf, J. Lin, R. Pang, L. B. Wei, L. L. Xu, F. G. Sheng, C. H. Li, G. J. Li, L. Y. Zhao and Y. L. Hou, Advanced Materials, 2014, 26, 4114-4120.
230 Z. W. Tay, D. W. Hensley, P. Chandrasekharan, B. Zheng and S. M. Conolly, Jee T Med Imaging, 2020, 39, 1724-1734.
229 W. Tong, H. Hui, W. Shang, Y. Zhang, F. Tian, Q. Ma, X. Yang, J. Tian and Y. Chen, Theranostics, 2021, 11, 506-521.
228 World Health Organization, https://gco.iarc.fr/daytoday/factsheets-cancers, (accessed May 2021).
227 A. Tomitaka, H. Arami, Z. Huang, A. Raymond, E. Rodriguez, Y. Cai, M. Febo, Y. Takemura and M. Nair, Nanoscale, 2017, 10, 184-194.
226 A. Antonelli, P. Szwargulski, E. Scarpa, F. Thieben, G. Cordula, G. Ambrosi, L. Guidi, P. Ludewig, T. Knopp and M. Magnani, Nanomedicine (Lond), 2020, 15, 739-753.
225 Y. L. Hou, H. Kondoh, T. Kogure and T. Ohta, Chemistry of Materials, 2004, 16, 5149-5152. DOI: 10.1021/CM035670K
224 Z. W. Tay, D. W. Hensley, P. Chandrasekharan, B. Zheng and S. M. Conolly, Jee T Med Imaging, 2020, 39, 1724-1734.
223 K. P. Melo, A. V. Makela, N. N. Knier, A. M. Hamilton and P. J. Foster, Magnetic Resonance in Medicine, 2021.
222 T. Knopp, S. Biederer, T. F. Sattel, M. Erbe and T. M. Buzug, Jee T Med Imaging, 2011, 30, 1284-1292.
221 S. B. Goldhaber, Regul Toxicol Pharmacol, 2003, 38, 232-242.
220 Z. W. Taylor, presented in part of World Molecular Imaging Congress (WMIC), Montreal, September 2019.
219 A. Antonelli, P. Szwargulski, E. Scarpa, F. Thieben, G. Cordula, G. Ambrosi, L. Guidi, P. Ludewig, T. Knopp and M. Magnani, Nanomedicine (Lond), 2020, 15, 739-753.
218 A. Tomitaka, H. Arami, Z. Huang, A. Raymond, E. Rodriguez, Y. Cai, M. Febo, Y. Takemura and M. Nair, Nanoscale, 2017, 10, 184-194.
217 World Health Organization, https://gco.iarc.fr/daytoday/factsheets-cancers, (accessed May 2021).
216 W. Tong, H. Hui, W. Shang, Y. Zhang, F. Tian, Q. Ma, X. Yang, J. Tian and Y. Chen, Theranostics, 2021, 11, 506-521.
215 A. Antonelli, P. Szwargulski, E. Scarpa, F. Thieben, G. Cordula, G. Ambrosi, L. Guidi, P. Ludewig, T. Knopp and M. Magnani, Nanomedicine (Lond), 2020, 15, 739-753.
C. Fink, M. Smith, O. C. Sehl, J. M. Gaudet, T. C. Meagher, N. A. Sheik, J. D. Dikeakos, M. J. Rieder, P. J. Foster and G. A. Dekaban, *Diagn. Interv. Imaging*, 2020, 101, 577-588.

R. Rohani, S. N. Chickera, C. Willert, Y. Chen, G. A. Dekaban and P. J. Foster, *Mol. Imaging Biol.*, 2011, 13, 679–694.

E. Obeid, R. Nanda, Y. X. Fu and O. I. Olopade, *Int. J. Oncol.*, 2013, 43, 5-12.

A. V. Makela, J. M. Gaudet, M. A. Schott, O. C. Sehl, C. H. Contag and P. J. Foster, *Mol. Imaging Biol.*, 2020, 22, 958-968.

A. V. Makela, J. M. Gaudet and P. J. Foster, *Sci. Rep.*, 2017, 7, 42109.

A. V. Makela and P. J. Foster, *Magn. Reson. Med.*, 2018, 80, 1138-1147.

M. Gerosa, G. Ren, Y. Zhang, P. W. Goodwill, J. Mansfield, P. Marzola and M. Wintermark, *Int. J. Mag. Part. Imag.*, 2020, 6, 2009029.

J. Mansfield, presented in part of World Molecular Imaging Congress (WMIC), Virtual, October 2020.

I. Baccelli, A. Schneeweiss, S. Riethdorf, A. Stenzinger, A. J. Mansfield, presented in part of World Molecular Imaging Congress (WMIC), Virtual, October 2020.

A. Rivera-Rodriguez, L. B. Hoang-Minh, A. Chiu-Lam, N. Sarna, S. Lefevre, D. Ruimy, F. Jehl, A. Neuville, P. Robert, C. Sordet, J. W. M. Bulte, P. Walczak, M. Janowski, K. M. Krishnan, H. B. Zheng, M. P. von See, E. Yu, B. Gunel, K. Lu, T. Vazin, D. V. Rybak, A. Schneeweiss, S. Riethdorf, A. Stenzinger, A. J. Mansfield, presented in part of World Molecular Imaging Congress (WMIC), Virtual, October 2020.

A. V. Makela, J. M. Gaudet, M. A. Schott, O. C. Sehl, C. H. Contag and P. J. Foster, *Mol. Imaging Biol.*, 2020, 22, 958-968.

A. V. Makela, J. M. Gaudet and P. J. Foster, *Sci. Rep.*, 2017, 7, 42109.

A. V. Makela and P. J. Foster, *Magn. Reson. Med.*, 2018, 80, 1138-1147.

M. Gerosa, G. Ren, Y. Zhang, P. W. Goodwill, J. Mansfield, P. Marzola and M. Wintermark, *Int. J. Magn. Part. Imag.*, 2020, 6, 2009029.

J. Mansfield, presented in part of World Molecular Imaging Congress (WMIC), Virtual, October 2020.

I. Baccelli, A. Schneeweiss, S. Riethdorf, A. Stenzinger, A. Schillert, V. Vogel, C. Klein, M. Saini, T. Bäuerle, M. Wallwiener, T. Holland-Letz, T. Höfner, M. Sprick, M. Scharpff, F. Marmé, H. P. Sinn, K. Pantel, W. Weichert and A. Trumpf, *Nat. Biotechnol.*, 2013, 31, 539–544.

A. Rivera-Rodriguez, L. B. Hoang-Minh, A. Chiu-Lam, N. Sarna, L. Marrero-Morales, D. A. Mitchell and C. M. Rinaldi-Ramos, *Nanotheranostics*, 2021, 5, 431-444.

S. Lefevre, D. Ruimy, F. Jehl, A. Neuville, P. Robert, C. Sordet, M. Ehlinger, J. L. Dietemann and G. Bierry, *Radiology*, 2011, 258, 722-728.

S. G. Ruehm, C. Corot, P. Vogt, S. Kolb and J. F. Debatin, *Circulation*, 2001, 103, 415-422.

D. B. Mangarova, J. Brangsch, A. Mohtashamidolatshahi, O. Kosch, H. Paysen, F. Wiekhorst, R. Klopfeisch, R. Buchholz, U. Karst, M. Taupitz, J. Schnorr, B. Hamm and M. R. Makowski, *Sci. Rep.*, 2020, 10, 12410.

W. Zakrzelewski, M. Dobrzynski, M. Szymonowicz and Z. Rybak, *STEM Cell Res. Ther.*, 2019, 10, 68.

M. F. Kircher, S. S. Gambhir and J. Grimm, *Nat. Rev. Clin. Oncol.*, 2011, 8, 677-688.

M. Zhang, X. Liu, J. Huang, L. Wang, H. Shen, Y. Luo, Z. Li, H. Zhang, Z. Deng and Z. Zhang, *Nanomedicine*, 2018, 14, 2475-2483.

M. Edmundson, N. T. Thanh and B. Song, *Theranostics*, 2013, 3, 573-582.

F. Chen, M. Ma, J. Wang, F. Wang, S. X. Chern, E. R. Zhao, A. Jhunjhunwala, S. Darmadi, H. Chen and J. V. Jokerst, *Nanoscale*, 2017, 9, 402-411.

M. Hofmann, K. C. Wollert, G. P. Meyer, A. Menke, L. Arseniev, B. Hertenstein, A. Ganser, W. H. Knapp and H. Drexler, *Circulation*, 2005, 111, 2198–2202.

F. Fidler, M. Steinke, A. Kraupner, C. Grüttnzer, K. H.iller, A. Briel, F. Westphal, H. Walles and P. M. Jakob, *IEEE Trans. Magn.*, 2015, 51, 5100704.

J. W. M. Bulte, P. Walczak, M. Janowski, K. M. Krishnan, H. Arami, A. Halkola, B. Gleich and J. Rahmer, *Tomography*, 2015, 1, 91-97.

K. Lüdtke-Buzug, D. H. Rapoport and D. Schneider, *AIP Conf. Proc.*, 2010, 1311, 244-248.

B. Zheng, M. P. von See, E. Yu, B. Gunel, K. Lu, T. Vazin, D. V. Schaffer, P. W. Goodwill and S. M. Conolly, *Theranostics*, 2016, 6, 291–301.
373 F. Lieb and T. Knopp, Medical Physics, 2021, 48, 3893-3903.
374 A. von Gladiss, M. Graeser, A. Behrends, X. Chen and T. M. Buzug, Sci. Rep., 2020, 10, 18432.
375 H. Bernd, E. De Kerviler, S. Gaillard and B. Bonnemain, Invest. Radiol., 2009, 44, 336-342.
376 D. T. Kehagias, A. D. Goulamios, V. Smyriiotis and L. J. Vlahos, J. Magn. Reson. Imaging, 2001, 14, 595-601.
377 A. Singh, T. Patel, J. Hertel, M. Bernardo, A. Kausz and L. Brenner, Am. J. Kidney Dis., 2008, 52, 907-915.
378 E. U. Saritas, P. W. Goodwill, G. Z. Zhang and S. M. Conolly, IEEE Trans. Med. Imaging, 2013, 32, 1600-1610.
379 I. Schmale, B. Gleich, J. Rahmer, C. Bontus, J. Schmidt and J. Borgert, IEEE Trans. Magn., 2015, 51, 6502604.
380 O. Dossel and J. Bohnert, Biomed. Tech. (Berl), 2013, 58, 611-621.
381 M. Graeser, P. Ludewig, P. Szwargulski, F. Foerger, T. Liebing, N. D. Forkert, F. Thieben, T. Magnus and T. Knopp, Physics in Medicine and Biology, 2020, 65.
382 J. Pagan, C. McDonough, T. Vo and A. Tonyushkin, IEEE Trans. Magn., 2021, 57.
383 E. E. Mason, C. Z. Cooley, S. F. Cauley, M. A. Griswold, S. M. Conolly and L. L. Wald, Int. J. Magn. Part. Imaging, 2017, 3.
384 M. Graeser, F. Thieben, P. Szwargulski, F. Werner, N. Gdaniec, M. Boberg, F. Griese, M. Moddel, P. Ludewig, D. van de Ven, O. M. Weber, O. Woywode, B. Gleich and T. Knopp, Nat. Commun., 2019, 10, 1936.
385 E. E. Mason, E. Mattingly, K. Herb, M. Sliwiak, S. Franconi, C. Z. Cooley, P. J. Slanetz and L. L. Wald, Sci Rep-Uk, 2021, 11.