Introduction

Magnetic Resonance Imaging (MRI) is a non-invasive imaging technique which forms images of the inside of the body [1,2]. MRI works on the principle of nuclear magnetic resonance (NMR), a phenomenon where nuclei of atoms get excited in the magnetic field by electromagnetic waves and emit signals [3,4]. Magnetic Resonance (MR) was independently developed by Felix Bloch and Edward Purcell in 1946. Bloch and Purcell shared the 1952 Nobel Prize in physics for this discovery [1,4]. Paul Lauterbur generated the first MRI on small test tube samples in 1973 [1]. Lauterbur and Mansfield formed different ways to generate images from magnetic spins [4]. They both were awarded the Nobel Prize in Medicine in 2003 for their discoveries about MRI [1].Since then, MRI has been an invaluable tool for diagnosis of a number of pathologies in medicine [2,5].

Basics of MRI

Electromagnetic waves travel at the speed of light ($3 \times 10^8$ m/sec). They form electric and magnetic field components, which are perpendicular to each other. X-Ray, visible light, microwaves, and radio waves are all forms of electromagnetic waves. In MRI, the 3 - 100 MHz range of radiofrequency (RF) pulse, an electromagnetic pulse, is used for signal generation. Bloch hypothesized that when any charged particle spins, it generates an electromagnetic field [3] (Figure 1). An atom may have an even or an odd number of protons. When there is an even number of protons, the paired protons cancel each other’s magnetic field resulting in a net zero magnetic field. With an odd number of protons, there is always an existence of one unpaired proton, which produces a net magnetic field or magnetic dipole moment (MDM) to the nucleus. While MRI could use any nuclei with an odd number of protons, hydrogen has been selected for imaging. The reason being that hydrogen is found in abundance in the human body.

Hydrogen is found in water, which is approximately 60% of the body weight, and in fats [3]. Protons of nuclei spin on their own axes and have their own small magnetic fields or MDMs. When there is no external magnetic field ($B_0$), the axes of these MDMs are randomly arranged. Thus, they cancel each other out resulting in a net zero magnetic field. In the presence of $B_0$, over time, the protons’ spins will line themselves up in the direction of $B_0$ and produce a net magnetic field. Protons start to wobble or precess in the presence of $B_0$. Thus, they spin on their own axes as well as precess about the axis of $B_0$. The rate of precession of protons around $B_0$ is based on the Larmor equation: $\omega = \gamma B_0$.

Abstract

Magnetic resonance imaging (MRI) is a powerful tool to get spatial information about muscles of different regions inside the body. It has the advantage of being non-invasive and being without any radiations. It is objective and quantitative tool. This review covers the basics of MRI and informs about how the image is obtained using MRI.

Abbreviations: MRI: Magnetic Resonance Imaging; NMR: Nuclear Magnetic Resonance; MR: Magnetic Resonance; RF: Radiofrequency; MDM: Magnetic Dipole Moment; FID: Free Induction Decay; PD: Proton Density
Where \( \omega \) = angular precessional frequency of a proton (expressed in MHz),
\[ \gamma \] = gyromagnetic ratio (expressed in MHz/Tesla),
\( B_0 \) = strength of the external magnetic field (expressed in Tesla) [3].

When the object to be imaged is placed in \( B_0 \), the proton spins align longitudinally along the direction of \( B_0 \) about which they precess. The magnetization along the direction of \( B_0 \) is called longitudinal magnetization, \( M_z \) [1,3]. When an RF pulse is applied, this longitudinal magnetization, \( M_z \) flips into a transverse plane, creating a transverse magnetization, \( M_{xy} \) (Figure 2) [2,4]. This transverse magnetization will precess around \( B_0 \) producing an alternating current in a receiver coil and thus, generating a measureable signal [2]. When the RF pulse is turned off, the protons start going back to their lower energy states, realigning in the direction of \( B_0 \). This causes a decay of magnetization over time, resulting in a decreased signal in the \( M_{xy} \) plane. This decreased amplitude of the signal over a period of time is called free induction decay (FID) (Figure 3) [2]. Two independent relaxation processes occur at the same time, namely, longitudinal relaxation and transverse relaxation [2].

**Relaxation Times**

When the protons’ spins go back to their lowest energy state or equilibrium state, it is called relaxation. Relaxation occurs when the RF pulse is turned off and protons realign along the direction of \( B_0 \), thereby releasing all the extra energy [3]. Relaxation time is the time required by the protons to go back to their lowest energy state [2].

**Longitudinal Relaxation Time:** Longitudinal relaxation time, \( T_1 \), is the time taken by the protons’ spins to realign themselves along the direction of the longitudinal (Z) axis [2,3,6]. \( T_1 \) is also called spin-lattice relaxation time because it also denotes the time required for the excited protons’ spins to give back the energy to the surrounding lattice in order to reach an equilibrium state [3,6]. Transverse relaxation time: Transverse relaxation time, \( T_2 \), is the time taken for decaying of transverse magnetization in the \( xy \) plane.

\( T_2 \) decay occurs 5 to 10 times faster than \( T_1 \) recovery (Figure 4) [3]. When the RF pulse is turned off, protons’ spins will start to dephase because of spin-spin interactions and external magnetic field inhomogeneities [2,3]. \( T_2 \) is also called spin-spin relaxation time because it is the time taken by transverse magnetization to decay which is caused due to spin-spin interactions. When the effect of external magnetic field inhomogeneities is added to \( T_2 \) relaxation time, it is called \( T_2^* \) relaxation time [7].

**Figure 4:** The longitudinal magnetization recovery and transverse magnetization decay occur independently of each other. \( T_1 \) is the time to regain 63% of original net magnetization. \( T_2 \) is the time of signal decay to 37% of its original signal [2].

**Figure 5:** RF pulse is given to the subject and then signal is obtained from magnetized spins in the body [1].
Image Formation with MRI: In the presence of B0, unpaired protons’ spins line up in the direction of the magnetic field. After the protons’ spins line up, an RF pulse is transmitted to the tissues of the body, causing some spins to align in the direction of a new magnetic field. After the RF pulse, magnetized spins in the body produce an MR signal while they return to their original alignment (Figure 5). But the signal obtained contains information from the entire imaged part of the body without any spatial information, so gradients are used to get spatial information. Gradients produce a linear magnetic field non-uniformity along all three axes which helps in obtaining spatial information [3].

There are three different types of gradients used [2-4]:

a) Slice-Selection Gradient – A slice-selection gradient is used to select a particular slice in the body [3]. For imaging a specific slice, an RF pulse of a specific frequency is chosen to only excite the selected slice with the slice-selection gradient in the z-axis [4].

b) Frequency-Encoding Gradient – Once the slice is selected, a frequency-encoding gradient is used along the x-axis after the RF pulse is applied. It affects the precession frequency of the protons, generating signals of different frequencies, which vary from each other in a predictable manner [4].

c) Phase-Encoding Gradient – A phase-encoding gradient is used for inducing phase differences along the y-axis. It is turned on and off for a short period of time, after the RF pulse is applied and before the application of a frequency-encoding gradient [4,6]. When this gradient is turned on, some protons start precessing faster than others, depending on their location in relation to the gradient. When the gradient is switched off, the protons become out of phase from one another. Thus, protons are in different phases along the y-axis [4].

Parameters that Define Contrast in an Image

Image Contrast: One of the most significant benefits of MRI, particularly when compared to other imaging techniques is its ability to differentiate between various tissues of the body [4,7]. Different signal intensities of different tissues produce an image contrast [8]. Intrinsic parameters of image contrast: T1 and T2 are fixed intrinsic properties of the tissue [3], which have already been described above. Contrast is generated between tissues because of differences in T1 and T2 values [7]. Another parameter which can define an image contrast is the proton density of the tissue [8].

Proton Density: Proton Density (PD) refers to the concentration of hydrogen protons in a tissue. It exists within the tissue and is thus independent of the existence of an external magnetic field. PD usually denotes the number of mobile hydrogen protons present in a tissue and is related to the signal intensity [9].

Extrinsic Parameters of Image Contrast: Image contrast can also be obtained by changing the operator-controlled parameters of MRI: repetition time and echo time.

Repetition Time: Repetition time (TR) is the time interval between two consecutive 90-degree RF pulses.

Echo Time: Echo time (TE) is the time interval between sending the RF pulse and measuring the signal [3]. By changing TR and TE, various “weighted” images can be formed. T1 weighted image: If TR and TE are both short, a T1 weighted image is obtained [3,10]. With a short TR, tissues can be distinguished from each other by their different T1s [3] (Figure 6). If TR is long, it would have caused a full recovery of longitudinal magnetization of tissues, resulting in no difference in their T1s. So, a short TR helps in increasing the T1 effect. A short TE results in reduction of the T2 effect [2,3,6,11,12]. A T1 weighted image provides good anatomical details of different tissues [6]. Fat has more signal intensity than water as recovery of fat is quicker than that of water [10]. Signal intensity is low for most lesions [10].

T2 Weighted Image: If TR and TE are both long, a T2 weighted image is obtained [3,9]. T1 effect is reduced when TR is long [2,3,12]. A TR of 2000-3000 msec can be used for decreasing the T1 effect (Hashemi, 2004). A long TE results in more dephasing of protons, which further increases the T2 contrast between tissues (Figure 7) [2,3,6]. Fat has less signal intensity than water as decay of fat is more rapid than that of water [9]. Signal intensity is high for most lesions [10].

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**Proton Density Weighted Image:** By using a long TR and a short TE, a proton density weighted image is produced. A long TR minimizes the T1 effect and a short TE decreases the T2 effect. So, the resultant contrast between the tissues would be based on the proton densities of the tissues [2,3,12].

**Conclusion**

Magnetic resonance imaging is an invaluable tool for getting the information about the inside of the body, in a safe way. It has been used for diagnostic and research purposes for a long time. The above review explains about the basic concepts so to have a better understanding about how MRI works.

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