Development of cardiac phantom which mimics the heart function using a 4.7t preclinical MRI system for cardiac imaging

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Abstract
Cardiac MRI was one of the first MRI applications in the 1980s, which helps in the human study of different cardiovascular diseases. MRI is a noninvasive imaging technique that can be used for diagnosis and has advantages over other imaging techniques since it does not rely on radiation like X-rays, PET, and SPECT. Modern clinical methods do not require much scanning time, it reveals good contrast between tissues and can compensate for patient movement during the scan. The scanner used is a 300mm internal diameter superconducting magnet (Magnex Uk) 4.7T preclinical MRI system, in the Biomedical Physics building at the University of Aberdeen. The cardiac phantom is made of a balloon filled with water, connected via a nylon T-piece to two 5 m lengths of rigid pvc tubing of 3 mm internal diameter to a 12 V water pump and solenoid valve located outside the 4.7 T MRI system’s 5 Gauss stray field contour. The balloon part of the phantom was fixed on the 4.7 T MRI system’s animal bed, directly over a 30mm diameter surface coil normally used to image the rat heart. The flow system was primed with water to mimic blood and the bed was loaded into the MRI magnet for imaging. The cardiac phantom has successfully been developed using a small balloon to represent a heart chamber and imaged in a 4.7T preclinical MRI scanner. Images acquired with gradient echo pulse sequence with shorter TE of 5 ms and using surface coil displayed improved image contrast and have useful signal-to-noise ratio as compared to images acquired with birdcage coil which show ghosting artifact.

Keywords: Cardiac phantom; Heart function; Cardiac imaging; Balloon phantom; 4.7T MRI; Heart chamber; Preclinical MRI.

1. Introduction
Cardiovascular diseases are one of the leading causes of death globally, the most common ones are atherosclerotic coronary artery disease and ischemic heart disease. Coronary artery disease leads to over 80% of sudden cardiac death, which occurs within 24 hours of the onset previous from patients with asymptomatic. Bartling, et al, (2010).

Myocardial Infarction (MI) also known as a heart attack occurs due to a prolonged lack of blood flow to the coronary muscle resulting in some cell death, depending on the time at which blood occluded. Thrombosis is a common cause of blood clotting, arising from coronary arterial narrowing by atherosclerotic. Coronary bypass surgery is one of the treatment methods for cases associated with myocardial infarction, using healthy vessels, and enzymes to break down clots Bartling, et al, (2010).

Animal models of human cardiovascular disease are mostly used since they are readily available and have aided the need to translate MRI (techniques) for clinical to preclinical laboratory and are useful in cardiac research Vallee, et al.
The theoretical background

Magnetic resonance imaging deals with the interaction of certain atomic nuclei in the body with an external magnetic field. The hydrogen atom is good for MRI since it is abundant in the human body in large quantities, most of it in water molecules, and is a single proton. Hydrogen nucleus spin is \( I = \frac{1}{2} \), which generates angular momentum \( p \), along the spin axis given by equation (1). Vaughan, et al, (2004)
\[ p = \hbar I \]  

(1)

Where \( \hbar \) is Planck's constant/\( 2\pi \).

\( B_0 \) is the symbol for the main magnetic field and is measured in Tesla (T) typical value range from 1.0 T to 3.0 T in clinical MRI, while preclinical MRI uses an even higher magnetic field of 4.7 T to 7 T. The quantum model of Nuclear Magnetic Resonance (NMR) states that only two orientations of the magnetic moment are allowed, in the presence of an applied magnetic field. The nuclear spin in an applied magnetic field has \( 2I + 1 = 2 \) possible orientations (Since \( I = 1/2 \)), either parallel or antiparallel to the direction of the field \( B_0 \), and this corresponds with the two energy states low and high, and with energy difference \( \Delta E \), given by Equation 2, *Vaughan, et al, (2004)*.

\[ \Delta E = \gamma \hbar B_0 \]  

(2)

\[ E = h\nu = \hbar \omega, \ h\omega = \Delta E = \gamma \hbar B_0 \]

(3)

where \( \nu \) is the resonance frequency for proton and \( \gamma \) is the gyromagnetic ratio.

At thermal equilibrium the spins are distributed between the high and lower energy levels, though there are more spins in the lower energy level than higher energy level. In a sample with large number of nuclei, the spins will align randomly between the high and lower energy levels. The equilibrium distribution between the higher and lower energy level is given by Boltzmann's Equation (4).

\[ \frac{N_L - N_H}{N_H} = \frac{\hbar \omega}{kT} = \frac{\gamma \hbar B_0}{kT} \]  

(4)

Where \( N_L \) is the number of spins in the lower energy level and \( N_H \) the number of spins in the higher energy level, \( k \) is Boltzmann’s constant and \( T \) absolute temperature.

Cardiac MR images are improved by the use of cardiac gating. During image acquisition, MR images suffer from motion artifacts caused by both cardiac and respiratory movement. But with modern imaging techniques, cardiac MR image qualities have improved by reducing the total time of image acquisition and by using cardiac and respiratory gating. Cardiac MRI is used to obtain the anatomical image of the heart and vessels, for patients with cardiac diseases to check for abnormalities. These images could be obtained at different views conventionally (axial, coronal, or sagittal views), but other views are carried out for most cardiac studies (long axis, short axis, or views of valves). The heart is imaged repeatedly at a single slice throughout the cardiac cycle, data collection takes place over multiple cardiac cycles *Bartling, et al, (2010)*.

Cardiac gating is used for MRI data acquisition of the cardiac cycle, which begins at the detection of a physiological event (R-wave of ECG serves as the trigger), to obtain an image slice at a particular time in the cardiac cycle. A typical acquisition might use a 128 x 128 pixel image matrix, so 128 ECG triggers collecting 1 phase encoding step at the triggering point *Eze, et al, (2018)*.

3. Materials and methods

The materials used for this work are; a 300mm internal diameter superconducting magnet (Magnex Uk) 4.7T preclinical MRI system, in the Biomedical Physics building, University of Aberdeen. Generates horizontal \( B_0 \) field, with actively shielded gradient coil (Magnex Uk) internal diameter 220mm, driven by linear amplifiers (7796, AE Tecron, USA), a birdcage RF coil was used to transmit and receive signal at Larmor frequency.

In order to generate a realistic flow and motion in the phantom, an external control unit was use to drive a pump with constant frequency to produce a flow rate and motion similar to values found in-vivo. The phantom used to represent the heart chamber is entirely non-magnetic and is placed in center of the scanner. While the pump, solenoid valve and control electronics circuit are kept at a distance from the system for safety. The system is placed at 1m high to level with the magnet isocentre from the ground. These was done to have a constant pressure at both ends of phantom. The phantom is made of balloon filled with water connected via a nylon T-piece of 5m length of rigid PVC (polyvinyl chloride) tubing of inner diameter of 3mm, connected to a 12V water pump and solenoid valve located outside the 4.7T scanner.
The waveform generated was designed with three NE555 database multivibrators. NE555 are precision timing circuits that are capable of producing accurate oscillation or time delays, it can operate with supplies of 5V to 15V and are capable of sourcing current up to 200mA.

An astable multivibrator; was used to set the flow pulse frequency. The frequency and duty cycle of the flow switching can be controlled independently. It is used to control the frequency range of 3.5 – 5Hz or 210 beats per minute. Connecting two external resistors and a single external capacitor, the frequency, and the duty cycle can be controlled independently as shown in equation(5).

\[
\text{Frequency} = \frac{1.44}{(R_A + 2R_B)C} \quad (5)
\]

Where expected frequency = 3.5Hz, \( R_A = 4.7\, \Omega \), \( R_B = 4.7\, \Omega \), \( C = 36\, \mu \text{F} \). The required capacitor is 36\( \mu \text{F} \), but the available capacitor is 33\( \mu \text{F} \), the output pin 3 of the NE555 is connected to a monostable multivibrator.

Monostable multivibrators were used to generate a pulse, with a duration independent of the astable oscillation frequency. And is triggered via a high pass filter connected to the output pin with a cut-off frequency of 3.4KHz. And is also used to set a gating duration. A gating trigger is required to enable the MRI system to generate gated images that align with the cardiac cycle. The R-wave of an ECG is the trigger.

Pulse Width Modulation (PWM), was used to control the duty cycle output from the astable multi-vibration operation which is fed into the output of the monostable operation (pin2) clock input. The duty cycle is controlled by the resistor value of \( R_A \) and \( R_B \)

\[
\text{Duty cycle (D)} = \frac{(R_A + R_B)}{(R_A + 2 \times R_B)} \quad (6)
\]

The output pin from NE555 monostable multivibrator is connected to an (IN4148 diode) which acts as a rectifier, and output from the diode to the resistors in series and to (2TX653)NPN power transistor which drives the solenoid valve. The transistor is connected in series with a current limiting resistor to a 12V voltage supply to control the solenoid valve, as shown in Figure 1.

![Figure 1: Flow control board](image)

The solenoid valve is controlled using a power transistor switching by multivibrator circuits which also generate a gating pulse to stimulate an R-wave trigger for gated MRI acquisitions.

The flow control circuits were soldered into fiber-glass strip-board as shown in Figure 1, for the three multivibrators with a power supply of 12 V from an external source. A diode is connected in series in the positive (+Ve) row to prevent reverse polarity and also a 220 \( \mu \text{F} \) capacitor between the power supply rows to limit motor noise.

3.1. The Flow Phantom Design (circulating system)

The balloon part of the phantom was fixed on the 4.7 T MRI system’s animal bed, as shown in Figure 3, directly over a 30mm diameter surface coil normally used to image the rat heart. The flow system was primed with water to mimic blood and the bed was loaded into the MRI magnet for imaging.
The heart chamber is modeled by a small rubber balloon fed with a pulsed water flow controlled by a solenoid valve and timing circuit.

**Figure 3** The heart chamber phantom mounted on the animal bed over a surface coil.

### 4. Results

**Figure 4** Output pulse from the astable multivibrator used to control the overall pulse rate (grey) and the high pass filtered output used to trigger subsequent monostable multivibrator stages (orange).
Figure 5 Output from monostable multivibrator use to control the solenoid valve drive (grey) and potential divider output used for MRI trigger (orange).

Figure 6 Gradient echo MRI scans of cardiac phantom acquired with transmit receive volume coil.
5. Discussion

Because of its ability to readily control the characteristics of the balloon, silicone gel was chosen for this work, its elastic modulus, and relaxation time of $T_1$ and $T_2$. The cardiac phantom as shown in Figure 2 above is made of a balloon filled with water connected through a nylon T-piece to two 5m lengths of rigid pvc tubing of 3mm internal diameter to a 12V water pump and solenoid valve located outside the 4.7T scanner.

The flow is switched on and off with the solenoid valve to cause the balloon to pulse as the pressure changes, simulating a beating heart chamber. The solenoid valve is controlled using a power transistor switching by multivibrator circuits which also generate a gating pulse to simulate an R-wave trigger for gating MRI acquisition.

Figure 4, is a voltage-time plot acquired with a digital storage oscilloscope (72-8395, Tenma, China) of the astable multivibrator output used to set the pulse rate in the phantom. The direct output, shown in orange has a period of approximately 200 ms, corresponding to a pulse rate of 5 Hz or 300 bpm. The high pass filtered output shown in blue is used to trigger the monostable pulse generators which drive the solenoid valve and MRI gating input. Figure 5 shows voltage-time plots of the trigger pulses from two monostable circuits acquired with a digital storage oscilloscope (72-8395, Tenma, China). The gray trace shows the output of the monostable used to control the solenoid valve drive, here, set for a pulse width of approximately 110 ms, and the orange trace shows the output of the MRI trigger monostable, with a width of approximately 0.57 ms and after attenuation by the potential divider of 4V.

From Figure 6 (A) image with motion, echo time (TE) = 15 mm shows artifacts along the phase encoding direction. (B) with motion and gating. TE = 15 mm. (C) acquired with motion and gating, and (D) acquired with reduced echo time TE = 11 mm. All images used repetition time (TR) = 200 ms, slice thickness 1 mm, the field of view FOV = 40 mm, 256 x 256 pixels, and bandwidth 50kHz.

Images acquired with reduced TE = 11 mm, (D) has useful SNR and reduced artifact compared to (C) with TE = 15 mm, same TR = 200 ms, and FOV = 40 mm.

The second set of images was acquired with surface coils since images in Figure 6A-D appear with much noise in the background signal and images in Figure 7 appear with reduced noise.

Figure 7 shows an image acquired with a surface coil with a higher signal-to-noise ratio (SNR), image (A) was acquired without motion, and image (B) was acquired with motion but introduces ghosting artifacts in the phase encoding direction. Slice thickness = 1, field of view (FOV) = 30mm, echo time (TE) = 10ms, repetition time (TR) = 500ms, matrix 256 x 256 pixel. Values of SNR measured at 4.7 T for images acquired with gradient echo MRI scan as shown in Figure 7, for A = 21170 ± 264 and B = 1251 ± 90, these figures being the mean ± standard deviation over the imaged signal.
6. Conclusion

The cardiac phantom has successfully been developed using a small balloon to represent a heart chamber and imaged in a 4.7T preclinical MRI scanner. Though, with high frequency, the pump makes a click sound and when the phantom is not at the same height as the control unit, there is a collapse in the pump cycle.

Images acquired with gradient echo pulse sequence with shorter TE of 5 ms and using surface coil displayed improved image contrast and have useful signal-to-noise ratio Fig7, compare to images acquired with birdcage coil which show ghosting artifact resulting from motion as seen in Figure 6a-d. Also has reduced image acquisition time and shows the myocardia movement through the image plane during the cardiac cycle.

In this research work, the cardiac phantom was imagined in air, but an improvement in this work would be, to place the phantom in a tube surrounded with water, to have a representative of the nuclear magnetic resonance (NMR) characteristic of a mouse myocardial function and its relaxation time ($T_1$, $T_2$ values), and flow velocity in the phantom. Or the use of a thicker-walled material with properties closer to those of cardiac muscle, like PVA (polyvinyl alcohol) cryogel.

This research work used a variable resistor to adjust the frequency and duty cycle of the pump, this could be replaced to have a constant frequency and duty cycles to avoid the need of changing the operational cycle of the pump and have a constant output.

Compliance with ethical standards

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Disclosure of conflict of interest

All authors are in the same area of interest. Biomedical.

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