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Développement d'une nouvelle méthode de mesure du rythme cardiaque et du débit sanguin fondée sur les perturbations localisées d'un champ magnétique.

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Résumé

L'objectif principal de cette thèse est la conception, la réalisation puis la mise en œuvre d'un dispositif de mesure pour l'acquisition du rythme cardiaque et du débit de circulation sanguine sans aucun contact ni électrique ni optique avec le patient, ces paramètres étant représentatifs physiologiquement pour évaluer la condition de santé d'un individu.

Ce travail s'appuiera sur une nouvelle méthode de mesure appelée MMSB (*Modulated Magnetic Signature of Blood*), fondée sur les perturbations localisées d'un champ magnétique rémanent au passage du flux sanguin. En effet, l'uniformité du champ magnétique généré par un aimant permanent de 0,1 – 0,2 Tesla placé à proximité de la peau d'un patient est perturbée par la circulation naturelle du sang suite aux propriétés ferromagnétiques des globules rouges. Cette perturbation est alors extraite sous la forme d'une tension électrique à l'aide d'un capteur GMR classique. L'originalité du dispositif repose sur une méthode d'acquisition non invasive et sans contact, permettant donc l'acquisition des signaux à travers les vêtements, la transpiration, les salissures ou autres polluants dans l'environnement proche du capteur, contrairement aux dispositifs commerciaux disponibles en début de thèse.

Un état de l'art détaillé des différentes techniques non-invasives liées à la mesure du rythme cardiaque (électrique, optique, micro-ondes, acoustique, mécanique et magnétique) est proposé en début de thèse, mettant en avant les avantages et inconvénients de chacune. Une présentation similaire est proposée pour comparer les méthodes connues et utilisées dans le cas de l'analyse du débit sanguin (dilution, thermo-dilution, par ultrasons...). Cette première partie permet de préciser ensuite les orientations et motivations vers une méthode de mesure magnétique.

Le manuscrit présente ensuite la méthode originale développée ici pour la mesure du rythme cardiaque et du flux sanguin de manière autonome (faible consommation), et non invasive, à savoir la méthode appelée MMSB. Une première étude permet la validation in vitro du phénomène MMSB avec la mise en œuvre d'un banc expérimental comprenant un aimant permanent couplé à un capteur GMR, un module d'acquisition et de traitement des signaux et un système de visualisation des données. Le dispositif d'étude in vitro est constitué d'un tube de verre rigide au travers duquel circule une solution d'oxyde de fer (Fe_2O_3) avec différentes concentrations correspondant à

quelques exemples possibles de quantité d'hémoglobine dans le sang parmi ceux rapportés dans la littérature. Cette étude se poursuit par une validation in vivo de l'exploitation du phénomène MMSB à l'aide d'une étude comparative de la forme d'onde obtenue à l'aide du capteur GMR et de l'amplitude MHV sur 20 sujets sains. L'analyse statistique des résultats montre une bonne concordance (erreur inférieure à 5%) sur les pics relatifs aux activités du cœur, les courbes de références étant obtenues avec une mesure étalon par ECG. Cette partie démontre ainsi les potentialités de la méthode MMSB proposée comme une alternative à l'ECG pour la mesure de la fréquence cardiaque chez les sujets sains, dans une situation de repos ou dans le cas d'environnements particuliers (ex : cas d'un corps dans l'eau tel que des nageurs ou plongeurs).

En parallèle, une étude multi physique est proposée pour étudier le phénomène MMSB. La construction d'un modèle est réalisée en deux étapes :

- modélisation bidimensionnelle du flux sanguin en présence d'un champ magnétique,
- modélisation du système d'écoulement fluide de l'oxyde de fer F_2O_3 dans un champ magnétique faible.

Des simulations sous le logiciel commercial multi physique COMSOL utilisent ce modèle, avec comme paramètres les valeurs relatives au banc expérimental ou des valeurs issues de la littérature. Ce modèle permet d'estimer la sensibilité de la méthode MMSB et d'optimiser la distance entre l'aimant permanent et le capteur. La comparaison entre les mesures et les résultats de simulation sous COMSOL montrent une bonne corrélation sur la valeur de cette distance optimale.

Pour aller plus loin, le phénomène MMSB est exploité non plus pour mesurer le rythme cardiaque, mais pour acquérir le débit sanguin en mesurant la variation de l'amplitude du signal après une période dite d'occlusion. En effet, l'amplitude du signal MMSB diminue pendant une occlusion correspondant à une chute du débit sanguin. Cette mesure a été comparée quantitativement et qualitativement en utilisant la technique de référence PORH (Post Occlusive Reactive Hyperaemia). Pour l'analyse qualitative, il s'agit d'évaluer le temps de transit entre la phase d'occlusion et le flux maximum détecté, qui est également mesurable par Laser Doppler. Une forte corrélation a été obtenue.

Pour l'analyse quantitative, une campagne de mesure sur 6 sujets sains montre une dispersion de moins de 4% entre la méthode de référence DRT4 et la méthode MMSB.

Ensuite, dans le cas de la surveillance continue de signes physiologiques tels que la pression artérielle, les travaux comparent l'application de la méthode MMSB pour l'acquisition du temps de transit impulsionnel (PPT) au niveau du poignet par rapport à l'acquisition au niveau du doigt à l'aide de photoplethysmogram (PPG) issues d'un appareil de référence OMRON.

Enfin, la description de l'implémentation matérielle est donnée, qui comprend l'électronique d'acquisition et de traitement, ainsi que l'encapsulation dans un dispositif portable faible consommation et autonome. Cette implémentation aboutit à la mise en œuvre d'une plate-forme de détection et d'analyse du rythme cardiaque à l'aide d'un réseau sans fil de type ad-hoc. Deux exemples d'application sont présentés, et deux dispositifs ont été entièrement développés lors de cette thèse et sont en cours de commercialisation, pour des applications de vie quotidienne, après avoir été préalablement brevetés.

Summary

The state of the art technologies and techniques to acquire heart or pulse rate and blood flow monitoring can be generally classified into electrical, optical, microwave, acoustic, mechanical and magnetic. The advantages and disadvantages for each these methods to acquire blood pulse and flow measurements will be discussed in this dissertation. Through these discussions, it was found that a novel method, capable of field deployability, is required to support the continuous monitoring of blood pulse and flow through fabric, body fluid and environment contaminants. Such a method must be small, capable of operations at low electrical power and is able to operate at room temperature. In addition, in order to support long period of usage, this method must also be able to operate without the need of a good electrical or optical contact.

This dissertation presents a novel and patented method of blood pulse and flow monitoring that is low power, small and capable of operations at room temperature. The method of blood pulse and flow monitoring is termed Modulated Magnetic Signature of Blood (MMSB) where a uniform magnetic field is applied on the skin, in a non-invasive manner, within close proximity of a major blood vessel and a Giant-Magneto-Resistance (GMR) based magnetic sensor of dimensions 6mm (length) x 5mm (width) x 1.8mm (height). The uniform magnetic field generated from a small permanent magnet, of dimensions 6mm (diameter) x 3mm (height), has a magnetic field strength of 0.1 - 0.2 Tesla. The applied magnetic field will create a uniform magnetic field encompassing the GMR based magnetic sensor, skin, fabric and blood vessel. The uniform magnetic field, when undisturbed, will produce a fixed DC output voltage from the magnetic sensor. However, due to the pulsatile nature of blood flow in the arteries, the uniform magnetic field will be disturbed periodically where the GMR based sensor will translate this magnetic disturbance as an output electrical voltage.

To demonstrate the MMSB phenomenon, an experimental setup was developed in this research using the flow of using iron (II) oxide (Fe_2O_3) fluid with air-gaps in a rigid glass tube. The sensor and magnet were placed at 2-3mm below the glass tube. Measurements were successfully conducted using different levels of Fe_2O_3 concentrations for each experiment, where each level Fe_2O_3 concentrations is selected based on published results on the amount of haemoglobin in blood. The results obtained from these experiments substantiated the MMSB phenomenon where measurements were subsequently conducted on human subjects successfully.

The MMSB phenomenon is thoroughly explored through physical measurements on locations where the major arteries are near to the skin. Each of these locations, such as the temple, neck, wrist and heel, was used for the verification of the MMSB phenomenon. Results obtained for measurements at all these locations were encouraging as the MMSB waveforms obtained were repeatable and are highly correlated to the activities of the heart.

The heart rate obtained using the MMSB phenomenon is quantitatively assessed with respect to the pulse and heart rate acquired by lifestyle (i.e. pulse oximeter) and clinical device (i.e. ECG measurement equipment) capable of supporting heart rate monitoring. The instantaneous heart or pulse rate obtained from the ECG and MMSB waveforms shows ± 2 *Standard Deviation (SD) of 0.48 blood pulse per minute (bpm), which is not significant for measurement of resting HR (typically 70 bpm) with variations of at least ± 1 bpm. Together with statistical tools used for validation of biomedical instruments, results presented in this dissertation support the use of MMSB as a viable alternative to ECG for measuring resting HR for healthy individuals.

The MMSB waveform was qualitatively analysed with respect to published results on activities of the heart. From these results, it was established that the peaks of the MMSB waveform have good correlations to the physical activities of the heart. As such, it is concluded that the MMSB phenomenon is qualitatively and quantitatively correlated to the physical and electrical heart activities; and is capable of reliable and accurate heart rate measurements.

Further to the physical measurements, the MMSB phenomenon is also modelled mathematically and simulated in a multi-physics simulation environment, COMSOL. Simulations for this model were successfully completed using parameters extracted from published information on blood.

Based on the results obtained from the model developed in COMSOL, it was concluded that the simulated waveform from the GMR sensor is a function of the pulsatile flow of blood. In order to compare the physical measurements with the simulated output, the distance between the magnet and the GMR sensor was varied in both physical measurements and simulation model. From these comparisons, it was established that the amplitudes are highly correlated and it was concluded the model is capable of modelling the MMSB measurements on human subjects.

To demonstrate the application of MMSB phenomenon for blood flow, a measurement setup was developed in this dissertation to correlate the changes on the amplitude of the MMSB pulse acquired on the wrist with respect to an occlusion on the upper limb. In this measurement, it was established that the amplitude of the MMSB pulse (both DC offset and pulsatile waveform) reduces

as the occlusion pressure increases. The direct correlation between the occlusion pressure and the amplitude of MMSB pulse substantiates the ability of MMSB in measuring blood flow.

The occlusion measurement using MMSB phenomenon was further developed for the post-occlusive reactive hyperaemia (PORH) measurements. The medical instrument capable of laser Doppler perfusion monitoring was used as the reference measurements. Qualitative and quantitative analysis were performed for the PORH measurements, using both MMSB and laser Doppler method, on six healthy subjects. Qualitative results obtained from these experiments demonstrated the ability of the MMSB phenomenon in acquiring the PORH waveform. Quantitative results obtained from the MMSB and laser Doppler method for PORH measurements were found to have errors of 4% or less. As such, the PORH measurement substantiates the capability of the MMSB phenomenon in both qualitative and quantitative blood flow measurements.

In addition, this dissertation established a MMSB experiment capable of delivering a feasibility study on using MMSB waveforms for the measurement of Pulse Transit Time (PTT). Using the measured pulse transit time and physical parameters obtained during the experiment, the Mean Arterial Pressure (MAP) of each subject can be calculated. For this experiment, Diastolic and Systolic blood pressure measured using commercial Automatic Blood Pressure (ABP) meters, MMSB and ECG data were collected for 10 subjects. These data were quantitatively assessed using the mean arterial pressure and were found to be correlated with less than 8% error. This error was found to be contributed by the uncertainties on the measurement parameters. With the exclusion of the uncertainties contributed by these measured parameters, it was mathematically estimated that the computed uncertainty can be reduced to 2%. As such, it was concluded that the use of MMSB and ECG data for PTT measurements, leading to the determination of the MAP data, has been successfully demonstrated.

Finally, in the application chapter of this dissertation, the physical implementation, including electronics design and packaging method, of the proposed MMSB method as a wearable device capable for supporting remote monitoring of blood pulse was presented. Together with the physical implementation, two applications of the MMSB phenomenon were also described. In both applications, the method of packaging applied is based on a second patent filed where the novelty in the method of packaging to achieve the robustness in deploying the MMSB phenomenon for end-user applications.

The first application is the funded development (\$350,000 by Singapore Totalisator Board) for a wearable, wrist based, wireless pulse sensing solution, where the MMSB phenomenon is integrated

with a low cost wireless ad-hoc network for continuous pulse rate monitoring. The system developed is capable of concurrent pulse rate monitoring of multiple users. This development had also won the SILVER award for the prestigious national level competition held in Singapore for innovative products and solutions. Further to the wireless pulse sensing, this development was enhanced with embedded processing to support localized display of measured heart rate on the Liquid Crystal Display (LCD) for health monitoring.

The second application of MMSB is the development of a lifestyle solution for measurement of changes in blood volume pulse for relaxation assessment. In this development, the MMSB phenomenon is embedded into an existing lifestyle solution capable of massaging the subject to make him/her relax. A statistical based algorithm was developed to extract the changes in blood volume pulse where the results were compared to the laser Doppler medical instrumentation for blood perfusion measurements. A trial on 40 subjects was conducted where the blood perfusion data was obtained before and after the massage using both the laser Doppler medical instrumentation and MMSB integrated product. The MMSB integrated product was able to produce measured results with 98% correlations to the laser Doppler measurements.

In addition, heart rate variability (HRV) was implemented as part of the algorithm to assess the well-being of the subject. The successful implementation and validation using statistical tools demonstrated the feasibility of using MMSB for HRV and is currently being developed as a lifestyle product.

Both applications described in the last chapter of the dissertation had attracted interest from companies and institutes in various countries such as Singapore (OSIM International, HuttCab Consulting and DSO), United States (JawBone and George Washington University), England (Swimnovate) and India (Amrita School of Engineering). At the point of writing this dissertation, development of commercialization prototypes has been completed for a Singapore based Multi-National Company (MNC) for a lifestyle product. At the same time, measurements were successfully duplicated in the laboratory of JawBone in California, United States and DSO in Singapore.

With the successful development of the novel sensing and packaging method using MMSB for blood pulse, flow and pressure analysis, this research has resulted in the filing of four patents and scientific publications of results in seven international conferences and journal. Research is on-going on new applications of MMSB for health management where future advances in the system architecture design, circuit design and signal processing can broaden the application potentials of physiological assessment for military, clinical or medical research. In addition, the unique capability of MMSB to

achieve continuous blood pulse, flow and pressure sensing through fabric, blood and light bandages can be exploited in future development to support the needs of commercial and medical research applications.

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Chapter 1: Introduction and Technology Overview

1 Introduction

With the advancement of bioelectronics, portable health monitoring devices are getting popular to provide continuous monitoring of an individual's health condition with ease of use and comfort. The portable health monitoring devices are increasingly used at places such as home, ambulance and hospital, and at situations military training and sports. Amongst the various physiological vital signs, heart or pulse rate and blood flow are two common parameters that are frequently monitored as they can be used to represent the normal operations of the heart. This chapter will present the state of the art technologies and techniques to acquire pulse rate and blood flow.

2 Pulse rate

Pulse rate is a measurement of the number of times the heart beats per minute. Typically, a normal resting heart rate in adults is 50 to 90 beats per minute [1] and the measurement of pulse amplitude and contour is a function of the pulsatile motion of the heart, which pushes blood through the arteries. This will result in the expansion and contraction of the blood vessels allowing blood to flow.

Pulse amplitude and contour, such as a small, weak pulse, a large rebounding pulse, or an irregularly shaped pulse has been reported to be indicative of a pathophysiologic state [1]. In addition, pulse diagnosis is also one of four kinds of diagnostic methods used in Traditional Chinese medicine (TCM) clinical practice to determine the physiological condition of patient [2]. Therefore, heart or pulse rate is an important physiological vital sign subject to continuous monitoring because they are representative in assessing the physical health condition of an individual. Healthcare institutes such as the hospitals and elderly care centres can use this information to monitor the health conditions of their patients. This is of particularly importance for patients with cardiac arrhythmia where heart rate variability needs to be monitored closely for early detection of cardiac complications.

Furthermore, pulse rate information of individuals subjected to mentally or physically stressful conditions may be utilized to trigger alert for immediate attention when large changes in heart rate variability potentially indicate fatal events such as heat stroke, cardiac disorder and mental break

down. Finally, it is also important to monitor pulse rate of personnel working in dangerous environments such as deep sea condition (divers), high temperature (fire-fighters), and deep underground (coal miners). Such monitoring together with other physiological vital signs, such as blood pressure and respiration rate, will minimize occupational risk and facilitate rescue operations in the event of any mishap.

2.1 Methods of non-invasive heart or pulse rate measurements

The use of electrical probes to measure heart rate (electrocardiogram or ECG) was discovered in 1872 by Alexander Birmick Muirhead. Alexander was reported to have attached wires to a feverish patient's wrist to obtain a record of the patient's heartbeat while studying for his Doctor of Science (in electricity) in 1872 at St Bartholomew's Hospital. This was directly recorded and visualized using a Lippmann capillary electrometer by the British physiologist John Burdon Sanderson. Following the successful recordings and visualization, Augustus Waller was the first to systematically approach the heart from an electrical point-of-view.

After the discovery of ECG, many methods of heart or pulse rate measurements had been developed. To-date, methods of heart or pulse rate acquisition can be classified into electrical[3],[4], optical [5],[6], microwave [7], acoustic [8],[10],[13], mechanical [9],[11] or magnetic [12],[14],[15] means.

2.1.1 Electrical method of heart rate acquisition

The commonly used electrical method of heart rate acquisition is the electrocardiogram (ECG or EKG). It is a recording of the electrical activity of the heart over time, usually in a noninvasive manner via skin electrodes. The name electrocardiogram is made of the following parts: electro, because it is related to electrical activity, cardio, Greek for heart, gram, a Greek root meaning "to write". Such a method requires good electrical contact with the human skin and the performance is subjective to human body fluids (e.g. blood, perspiration, skin-oil) and environmental contaminants (e.g. mud, water, etc).

In 1901, Willem Einthoven invented the string galvanometer and assigned the letters P, Q, R, S and T to the various deflections of the ECG as shown in Figure 1. Using these assigned letters, he was able to describe the electrocardiographic features of a number of cardiovascular disorders by associating the different waveforms for each of the specialized cells found in the heart with the ECG signal as shown in Figure 2. Since then, many researches had been conducted and to-date, ECG

measurements can range from 3 to 12¹ electrodes depending on the measurement requirements. In all of these measurements, one electrode is used as the reference electrode to ensure such a system is less susceptible to electrical noise [4]. Recent research [3] has also reported a new method of ECG acquisition using contact free capacitive sensing. However, this method has been reported to be highly subjective to noise and motion artifacts.

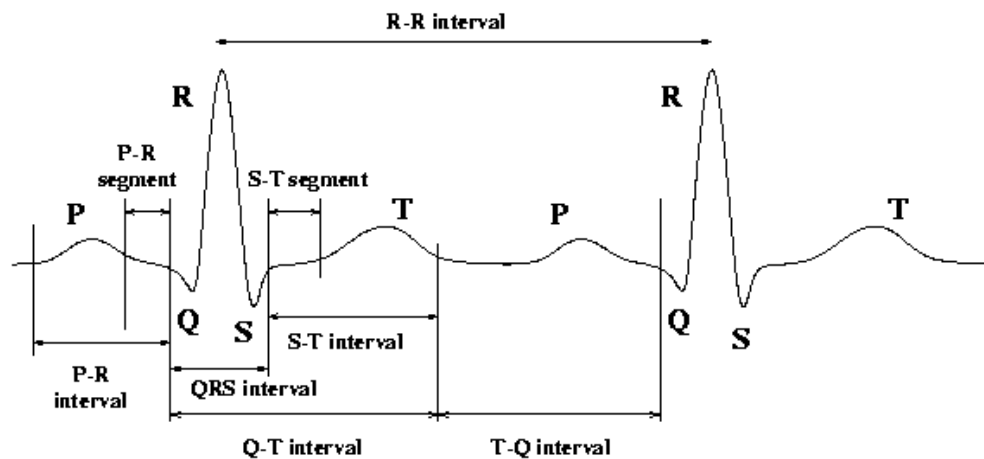


Figure 1 Definition of waves, segments and intervals in the normal ECG waveform

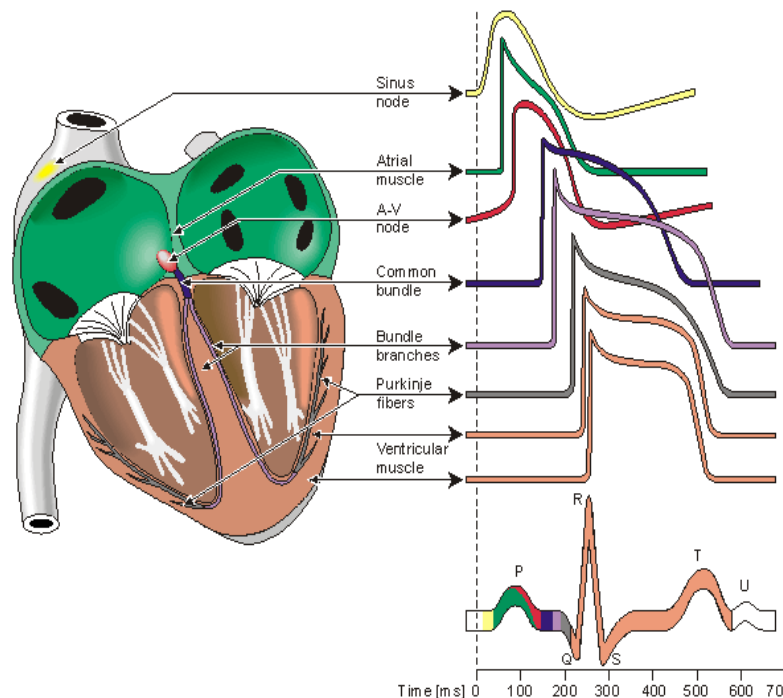


Figure 2 Association of different waveforms for each of the specialized cells found in the heart

Examples of commercially available heart rate acquisition systems include ECG monitoring devices, pulse rate monitoring watches and chest strips. All of these devices require electrical contact between the probes and the human skin and are usually subjective to motion artifacts.

¹ The 12 electrodes ECG is the gold standard for cardiac monitoring [16]

2.1.2 Optical method of heart rate acquisition

Pulse oximetry² was developed in 1972, by Aoyagi at Nihon Kohden using the ratio of red to infrared light absorption of pulsating components at the measuring site. It was commercialized by Biox in 1981 and Nellcor in 1983. Since then, the use of optical sensors in pulse oximetry [5],[6] is getting popular due to its compact nature and the ability to concurrently acquire SpO₂ concentration in blood and pulse rate. The basic operating principle of such a system requires a light source and a sensor with signal processing as shown in Figure 3. There are two methods to measure the light content where one method measures the reflected light while another measures the transmitted light. Both methods of measurements are able to determine the pulse rate and SpO₂ concentration in blood as a change in oxygen concentration in the blood will result in a change in light permittivity. Through the measurement of the change in light permittivity, the pulse rate and SpO₂ concentration in blood can be determined.

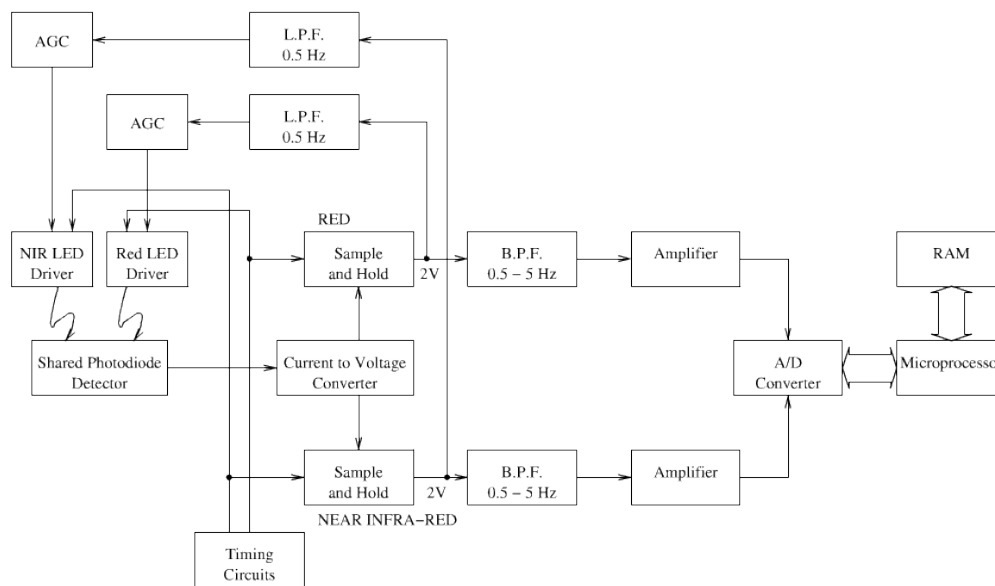


Figure 3 Block diagram of a typical pulse oximeter implementation

As the optical transmittance of the human skin is important to ensure signal quality, such a device is typically worn on the tip of the fingers or toes or on the earlobes allowing both the light transmittance and reflection methods to work.

To-date all of these devices require optical contact between the optical probes and the human skin. Motion artifacts are usually the cause of measurement noise and numerous algorithms had been developed to reduce such noise.

² Pulse oximetry is a non-invasive method allowing the monitoring of the oxygenation of a patient's hemoglobin through optical means.

2.1.3 *Microwave method of heart rate acquisition*

Microwave Doppler radar has been used to sense physiological movement since the early 1970s [17]. The original equipment was bulky, heavy and expensive. However, recent advances in micro-fabrication and wireless technology have led to the development of compact, lightweight and inexpensive microwave Doppler radar for measurements of physiological signs. The physiological signs are usually presented to the Doppler radar as periodic motion, such as that resulting from respiration and cardiopulmonary activities. In such systems, motion is measured as phase modulation and measurement sensitivity is significantly affected by phase noise and amplitude modulation sidebands in the local oscillator. Experimental results of heart rate measurements [6] under various signal output power conditions, with consideration for the impact of respiration had been reported for non-invasive detection of heart rate. However, typical systems based on the Doppler radar method have been reported to be subjective to motion artifacts, clothing and human body fluids [17].

2.1.4 *Acoustic method of heart rate acquisition*

The acoustic method of acquiring heart sound using an invention, called stethoscope, on the human chest was invented in France in 1816 by René-Théophile-Hyacinthe Laennec at the Necker-Enfants Malades Hospital in Paris. The stethoscope is an acoustic medical device for auscultation, or listening to the internal sounds of an animal body. It acquires the heart sounds, which are noises generated by the beating of the heart and the resultant blood flowing through the valves. In healthy adults, there are normally two heart sounds often described as a lub and a dub (or dup), that occur in sequence with each heartbeat. These are the first heart sound (S_1) and second heart sound (S_2), produced by the turbulent flow against the closed atrioventricular (AV) valves and semilunar valves of the heart respectively. Over the years, the stethoscope was progressively improved by Arthur Leared (1851), George Cammann (1852) and finally, Rappaport and Sprague designed a new stethoscope in the 1940s which became the standard by which other stethoscopes are subsequently developed.

In the recent years, scientist integrated electronics with the stethoscope and the electronic stethoscope (or stethophone) was created to overcome the low sound levels by electronically amplifying body sounds. However, amplification of stethoscope contact artifacts, and component cutoffs (frequency response thresholds of electronic stethoscope microphones, pre-amps, amps, and speakers) limit electronically amplified stethoscopes' overall utility by amplifying mid-range sounds, while simultaneously attenuating high- and low- frequency range sounds [17]. Through the years,

innovative signal acquisition and processing methods were developed to improve on the quality of the signal acquired [8],[10],[13] and the stethophone is now available with wireless and storage capabilities [17]. However, typical stethoscope or stethophone are known to be subjective to noise due to motion artifacts, clothing and environmental auditory noise.

2.1.5 Mechanical method of heart rate acquisition

Mechanical methods to acquire heart or pulse rate varies from the use of pressure cuff to piezo-electric materials worn over the limbs or body [9],[11]. Typical mechanical methods to measure pulse rate can be applied on blood pressure equipment using the oscillometric methods. This method comprises of an inflatable (Riva-Rocci) cuff placed around the upper arm at roughly the same vertical height as the heart. An electronic pressure sensor (transducer) is fitted in the cuff to detect blood flow. The cuff is inflated to a pressure initially in excess of the systolic arterial pressure, and then reduces to below diastolic pressure over a period of about 30 seconds. When blood flow is blocked (cuff pressure exceeding systolic pressure) or unimpeded (cuff pressure below diastolic pressure), cuff pressure will be essentially constant. Typically, both the blood pressure and pulse rate are acquired during this period. However, such mechanical methods to acquire heart or pulse rate requires the application of localized pressure through the cuff on the human subject and are not well suited for continuous signal acquisition.

2.1.6 Magnetic method of heart rate acquisition

Biomagnetic signal from the heart was detected in 1963 by Baule and McFee. The basic principle involves the mapping of the magnetic field around the thorax while the heart magnetic vector is acquired and is commonly known as magnetocardiogram (MCG). Typically, such a method requires highly sensitive magnetic sensors, such as SQUID (Superconducting QUantum Interference Device), and a magnetically shielded room to ensure minimum magnetic disturbance, including earth magnetic field. As such, SQUID is usually deployed in hospital while ECG is commonly found in clinical applications as it is proven to be more reliable, convenient and less expensive. However, there are still many potential clinical applications of the MCG. For instance, based on published data [12], with the combined use of the ECG and the MCG, called electromagnetocardiogram, (EMCG), the number of incorrectly diagnosed patients, for some cardiac diseases, can be decreased by one half of that when only the ECG is used.

The pulse rate can also be obtained by applying a magnetic field on the human blood and measuring using induced voltage using the Hall Effect [12],[15]. Such system applies a uni- or bi-directional

magnetic field on the human body to create polarization of blood molecules. Electrodes are placed on the human skin near the applied magnetic field to pick up the potential difference (i.e. Voltage) created by the polarized blood molecules. Such method, when applied invasively can also be used to measure blood flow during clinical surgery. When used in a non-invasive manner, this method requires good electrical contact between the skin and the electrodes and therefore, it is not a common method used for pulse rate acquisition.

2.2 Summary on methods of non-invasive heart or pulse rate acquisition

Each of the methods for heart or pulse rate acquisition outlined in this section has its strength and limitations. The strength and limitations are dependent on the applications. Based on the application of the methods listed in this section for lifestyle products, a comparison table is shown in Table 1.

Based on Table 1, it can be observed that all methods of heart or pulse rate acquisition listed are subjective to motion artifacts. In addition, all methods, except mechanical, are able to support continuous measurements of pulse rate acquisition. However, the magnetic method is the only method that is able to acquire heart or pulse rate through fabric, body fluid and environmental contaminants. However, this method requires a special room and it is bulky, heavy and expensive. As such, this method is only used for clinical research and in hospital.

	Electrical	Optical	Microwave	Acoustic	Mechanical	Magnetic (SQUID)
Electrical contact with human skin	X					
Optical access to blood vessel		X				
Motion artefacts	X	X	X	X	X	X
Relatively higher power consumption		X	X			X
Large (not wearable)			X			X
Signal dependent on electrical noise	X		X	X		
Able to measure through fabric, body fluid and environmental contaminants						X
Continuous measurements	X	X	X	X		X

Table 1 Comparison on existing well established methods of heart or pulse rate acquisition

2.3 Motivation of a non-invasive pulse rate acquisition using magnetic means

The capability of acquiring pulse rate through fabric, body fluid and environmental contaminants is a differentiating feature for lifestyle products. As such, the magnetic method of pulse rate acquisition motivated my research on a compact, lightweight and inexpensive means to acquire blood pulse.

3 Blood flow

Blood flow is the flow of blood in the cardiovascular system and is typically measured by observing blood flowing through a section of the blood circulatory system. It is a physiological vital sign that is dependent on the heart pushing blood into the arteries and the resistance presence in the arteries, which will expand and contract allowing blood to flow [18]. Mathematically, blood flow is described by Darcy's law (which can be viewed as the fluid equivalent of Ohm's law) and approximately by Hagen-Poiseuille equation [19]. Details of the mathematical model for blood flow will be presented in Chapter 3 of this dissertation.

Blood flow is an important physiological vital sign subject to continuous monitoring because it is representative in assessing the physical health condition of an individual. It is an important parameter to monitor for individuals subjected to crowded and cramped conditions with limited physical activity where an alert may be triggered for immediate attention when blood flow anomalies, such as deep vein thrombosis, are detected. For paraplegic patients, it is important for blood flow anomalies to be detected early to avoid medical complications, such as pressure sores (decubitus) and thrombosis, as a result of limited decreased movement. In addition, monitoring of blood flow anomalies for patients after major surgeries is important to ensure smooth recovery.

3.1 Methods of blood flow measurements

Circulatory flow of blood originates due to the muscle contraction of the heart as illustrated in Figure 4. Since blood is an incompressible fluid, the conservation of mass (volume) in a closed hydraulic system is known as the principle of continuity. The stroke output of the heart is the principal determinant of circulatory blood flow and as illustrated in Figure 4, the stroke output of the heart occurs when the ventricles contract and blood is pumped to the lungs and body. The measurement parameter for the stroke output of the heart is the Stroke Volume (SV). SV is the volume of blood pumped by the right or left ventricle of the heart in one contraction. Specifically, it is the volume of blood ejected from the right or left ventricle during systole and the value is typically obtained by subtracting the end-systolic volume (ESV) from end-diastolic volume (EDV) for the right or left ventricle respectively [20].

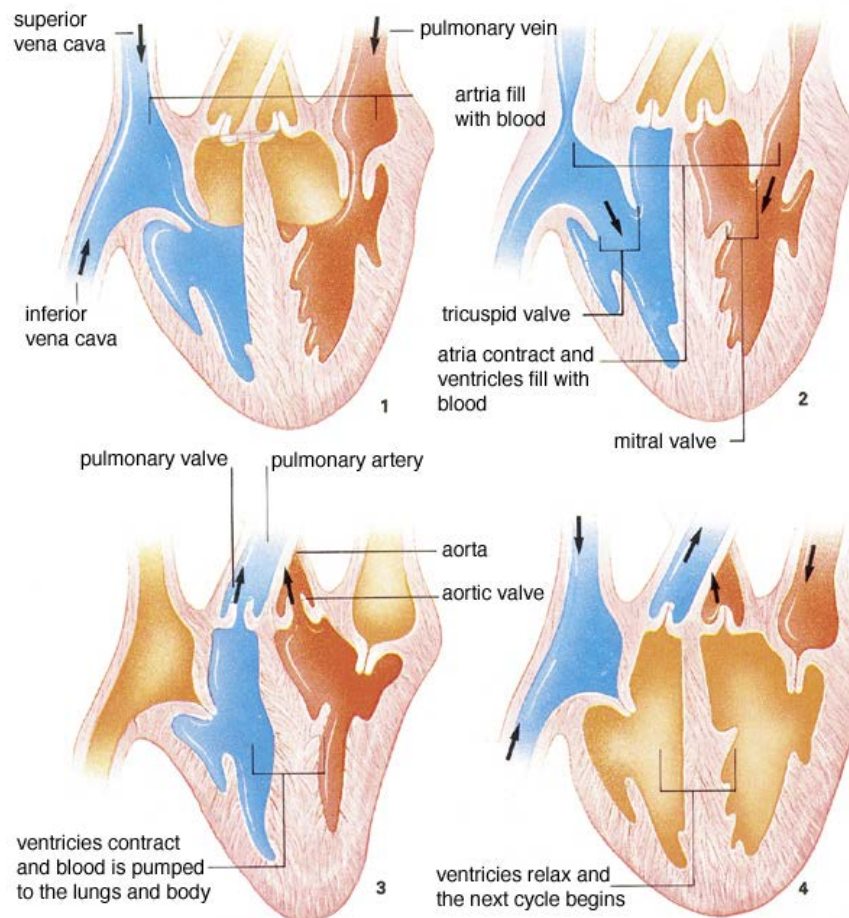


Figure 4 Illustration on operations of heart and blood flow during each step

On average, men have higher SV compared to women due to the larger size of their hearts. However, SV depends on several factors such as heart size, contractility, duration of contraction, preload (end-diastolic volume), and after-load. Typically, for most people, prolonged aerobic exercise training may increase stroke volume, which will, usually, results in a lower (resting) heart rate. This is because a reduction in heart rate prolongs ventricular diastole (filling), increases end-diastolic volume, and ultimately allows more blood to be ejected.

To quantify the volume of blood being pumped by the heart (ventricle) in a minute, the parameter used is the cardiac output, measured in $\text{L}\cdot\text{min}^{-1}$. Typically, an average cardiac output for human male is $5 \text{ L}\cdot\text{min}^{-1}$ and for human female is $4.5 \text{ L}\cdot\text{min}^{-1}$ [21].

To-date, there are a number of clinical methods for the measurement of the cardiac output. These methods range from invasive measurement using direct intra-cardiac catheterization to non-invasive measurement of the arterial pulse. Each method has its strengths and weaknesses where relative comparison is limited by the absence of a widely accepted “gold standard”. Therefore, the following sections will discuss a few of the specialized techniques developed to measure blood flow and

changes in blood volume. Each of these techniques differs in the measurement aspects of blood flow such as: the average flow; the instantaneous pulsatile flow; the flow profile; and changes in blood volume. These measurements are all unique and provide valuable information to physicians and medical researchers in clinical and research applications.

3.1.1 Indicator-dilution Method

Indicator-dilution method measures the averaged flow of blood in a vessel over a number of heart beats invasively. This method applies the concept of indicator, such as colored dyes and radioisotope, insertion into the blood stream and allowing it to flow in the blood circulatory system. Typically, the rate at which blood carries the indicator downstream is measured and two types of indicator-dilution methods that are clinically practice are: continuous infusion and rapid injection.

3.1.1.1 Continuous infusion techniques

The continuous infusion technique involves the continuous addition of an indicator into a flowing stream. For a fixed change in indicator concentration, ΔC ; at a known rate of injection, $\frac{dm}{dt}$; the average flow, F , can be calculated by Equation 1.

Equation 1
$$F = \frac{dV}{dt} = \frac{\frac{dm}{dt}}{\Delta C}$$

where V = volume of injectate (ml)

Many different indicators have been tried but the most successful indicator has been the indocyanine green, whose concentration curve can be measured in blood by light absorption. However, the main disadvantage of this method is the recirculation of most indicators where the indicator flows through the blood circulatory system and returns back to the injection point, affecting the maximum values of the cardiac output [22].

3.1.1.2 Rapid injection

In clinical use, continuous infusion has largely been replaced by rapid injection due to its clinical simplicity and reliability. Rapid or bolus injection method typically involves a bolus of indicator, such as a dye or radioisotope, being rapidly injected into the large vein or preferably into the right side of the heart [22]. The variation in downstream concentration of indicator, $C(t)$, is measured until the bolus has passed. The average flow is then obtained by Equation 2.

Equation 2
$$F = \frac{m}{\int_0^{t_1} C(t) dt}$$

where m = quantity of indicator injected

t_1 = time after which the indicator has completely passed.

3.1.2 *Thermo-dilution Method*

The invasive measurement method for cardiac output using temperature changes is thermo-dilution [22]. A bolus of cold saline is rapidly injected into the right or left atrium. The blood and saline mix in the right ventricle and the temperature difference is recorded in the pulmonary artery using a Thermistor. The Swan-Ganz Catheter is a four-lumen catheter clinically used for this procedure. It can stay within the patient's body for up to 24 hours; this allows a series of cardiac output measurements to be recorded based on the Equation 3.

Equation 3
$$\text{Cardiac Output} \propto (\text{blood temp} - \text{injectate temp})$$

3.1.3 *Electromagnetic flow meters*

The electromagnetic blood flow meter is an important tool with both medical research and clinical applications. It is based on the fact that blood is a conductor of electricity, so that the imposition of a magnetic field produces a difference of voltage across its diameter. Measuring this voltage using electrodes on the walls of the blood vessel will allow the measurement of the mean blood velocity or the instantaneous pulsatile flow of blood. The pulsation can be averaged with respect to time indicating average blood flow over a number of heart beats.

Typical electromagnetic flow meters provide measurement output for both the instantaneous pulsatile flow of blood and the average flow. For the instantaneous pulsatile flow of blood, the flow meter measures the induced electromagnetic field, e , depending on the fact that blood is flowing with velocity, u , through a magnetic field, B . This is given by Equation 4 [23].

Equation 4
$$e = \int_0^{t_1} (\mathbf{u} \times \mathbf{B}) dL$$

where L is the length between the recording electrodes.

For a uniform magnetic field and uniform velocity profile, the induced electromagnetic field is $e = B \times L \times u$.

In the case of parabolic flow profiles, resulting from laminar flow in tubes, the flow velocity u is replaced by the cross-sectional average flow velocity u_A . The volumetric flow can thus be attained by $A \times u_A$, where A is the cross-sectional area of the vessel.

There are both DC and AC flow meters where a comparison is tabulated in Table 2. Generally, due to the ability to differentiate signal from electrode offset potential and better noise immunity due to contact between electrode and tissues, the AC flow meters are commonly used in clinical application.

Flow Meter	Ability to differentiate signal from electrode offset potential	Noise due to amplifier drift	Immunity to noise due to contact between electrode and tissues
DC		x	
AC	x		x

Table 2 Comparison of DC and AC Blood flow meter [22]

AC flow meters are typically driven by sine or square waves, and use a toroidal-shaped cuff probe to create the magnetic field and properly position the electrodes. The probe must fit snugly around the blood vessel during diastole in order to record a valid signal. Therefore, as the vessel is slightly constricted during systole, this method is not applicable to veins as illustrated in Figure 5.

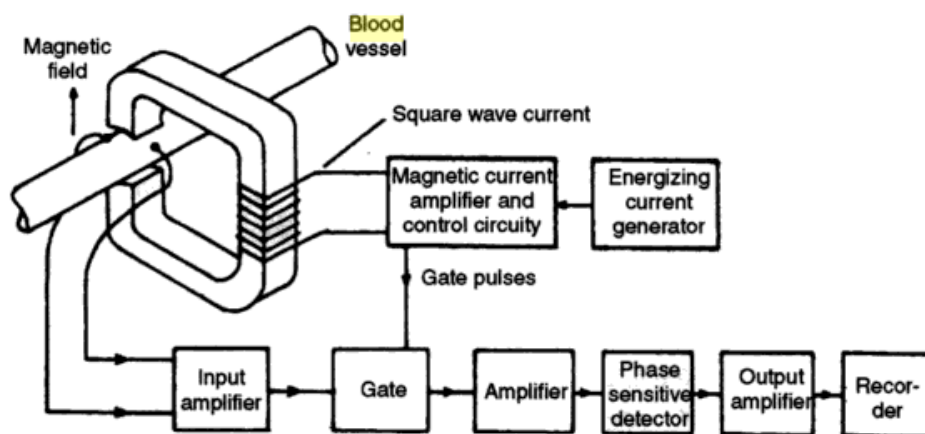


Figure 5 Typical block diagram of a square wave electromagnetic blood-flow meter [22]

3.1.4 Ultrasonic flow meters

The ultrasonic flow meter measures the blood flow in the ascending aorta by the application of Doppler principle (Huntsman et. al, 1983). It measures the instantaneous pulsatile flow in vessels non-invasively where advanced version of these devices, using Pulsed Doppler, can also measure flow profiles. The Pulsed Doppler flow meter applies an ultrasonic transmitter, which exerts a pulse travelling in a single packet to the source, i.e. red blood cell (RBC). The wave reflects off of RBCs in the blood stream and is received by the transmitter with a time delay proportional to the distance

travelled. Analyzing the Doppler shift at various delays creates the velocity profile of blood across the vessel.

Pulses are usually transmitted at 4 or 8 MHz and last about 1us [24]. The intensity of the packet is convolved with the velocity profile to create the reflected signal. Thus, the received signal must be mathematically de-convolved before it adequately represents the velocity profile across the vessel.

Since the introduction of the ultrasound Doppler instrument by M/s Deltex Medical, UK, many similar clinical based ultrasound Doppler instrument had been introduced and to-date, ultrasonic flow meters are widely used in clinical diagnosis. However, such method has not been widely adopted as a portable system for daily monitoring due to cost, size and complexity in operating the device, which requires a trained clinician.

3.1.5 Plethysmography

Impedance plethysmography, also called impedance test or blood flow or impedance phlebography, is a non-invasive test that uses electrical monitoring in the form of resistance (impedance) changes to measure blood flow in veins of the leg [25]. Impedance plethysmography uses the simple principle that electrodes placed on a tissue will be able to measure the resulting impedance of the tissue as shown in Figure 6. As the volume of the tissue changes, the resistivity changes leading to the impedance changes. A typical change in blood volume, ΔV , causes an increase in cross-sectional area, and thus leading to an increase in impedance, ΔZ . The relation for determining the change in blood volume is given by Equation 5 [26].

Equation 5
$$\Delta V = \frac{\rho_b L^2 \Delta Z}{Z_0^2}$$

where ρ_b = resistivity of blood³

L = distance between electrodes

Z_0 = impedance of the muscle tissues⁴

³ The resistivity of human blood is reported to be 1.6 Ωm [29]

⁴ The resistivity of muscle tissue is reported to be 3 Ωm [29] where impedance Z_0 can be estimated

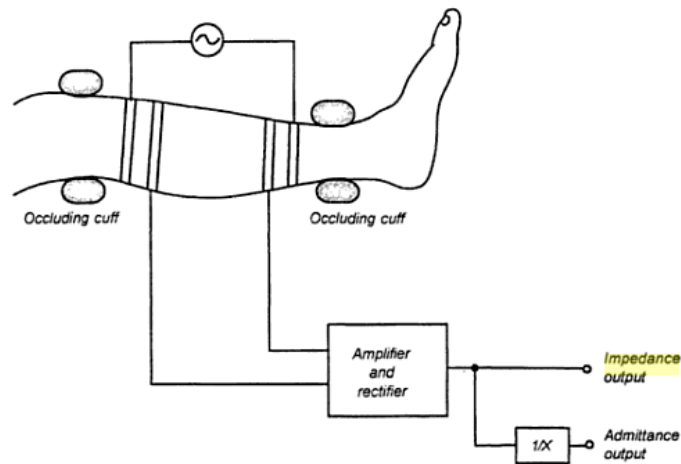


Figure 6 Impedance plethysmography system for limb blood flow measurement

Typical implementation of impedance plethysmography uses electrical probes to measure blood flow, known as Electric-impedance plethysmography. Such an implementation is non-invasive and the technique for measuring changes in blood volume is relatively simple as the electrodes can be placed at the any desired location on the body. For example, electrodes placed on limbs can detect vessel obstructions, and when placed around the neck and waist can detect beat-by-beat changes in cardiac output. However, the electric-impedance plethysmography is typically applied in the measurement of blood flow changes and is not commonly used for cardiac output measurement. As such, the electric-impedance plethysmography is a clinically accepted method to provide Information in helping doctors to detect deep vein thrombosis (blood clots or thrombophlebitis) [27].

3.2 Summary on methods of non-invasive blood flow signal acquisition

Most of the instruments for blood flow measurements are based on the sensing of velocity of blood instead of volumetric rate of blood flow. Based on the findings reported in [28], the electromagnetic and Doppler flow meter methods were compared with the thermo-dilution technique for the measurement of cardiac output in unanaesthetized rabbits. Cardiac output was varied by hemorrhage and transfusion and all three methods gave a satisfactory estimate of cardiac output over a wide range of controlled blood flows. The electromagnetic method is found to be the most accurate and the Doppler flow meter gives a systematic 3–4% overestimate of cardiac output. The thermo-dilution technique gives a good estimate of cardiac output at low and normal resting cardiac outputs, but is subject to a small but increasing error as cardiac output rises. The error at high flows is found to be of the order of 10%. In another report [30], comparison of blood flow measured by plethysmography and electromagnetic flow meter during steady state forearm exercise was done. It

was found that the plethysmography method tends to overestimate (up to 1.7 times) the blood flow of the subject in this experiment.

All these methods are either invasive or requires good electrical or optical contact with the skin limiting its deployment as a portable device for blood flow measurements. Therefore, to apply blood flow measurements beyond clinical deployment, research in blood flow signal acquisitions and analysis have been on-going to enhance these methods for a portable solution.

3.3 Motivation of a non-invasive blood flow signal acquisition using magnetic means

The potential application of the portable magnetic method for blood flow acquisition motivated the research of a compact, lightweight and inexpensive means to acquire blood flow. In order to achieve this, a mathematical model of the magnetic method of blood flow acquisition is derived, consisting of nonlinear partial differential equations obtained based on the application of Maxwell's Equation. The modelled equations are solved in a multiphysics simulation environment, which enables reliable predictive simulation of the underlying physical phenomena. The simulation results are compared with the corresponding experimental results and checked on their physical soundness.

In addition, research and development work presented in this dissertation will also outline experimental setups developed to demonstrate the application of the magnetic method for blood flow acquisition in a small and robust form. These experiments aim to provide the proof-of-concept for new lifestyle and medical products capable of non-invasive acquisition of blood flow signal through fabric, body fluid and environmental contaminants.

4 Conclusions

A technology review is presented in this chapter highlighting the advantages and disadvantages of each method used for blood pulse and flow measurements. It can be concluded that there is a lack of a portable system capable of measuring blood pulse and flow through fabric, body fluid and environment contaminants. Therefore, the motivation of this research is to develop a novel method of blood pulse and flow acquisition that is capable of deployment, in a robust manner, as a portable physiological monitoring system. Such a method will support the acquisition of both blood pulse and blood flow without the need for a good electrical or optical contact and can be used over a prolonged period of time on the limbs.

As such, one of the objectives of this project will be the development of a non-invasive magnetic based blood flow acquisition system that applies commercially available magnetic sensors in place of SQUID or electrodes. This will allow the system to operate at room temperature, be reliable, compact (small size), cheap, and convenient for daily usage.

Using this novel method to detect blood pulse and blood flow, systems can be developed for applications such as lifestyle or health monitoring device at home, ambulance and hospital, and at situations, including military training and sports.

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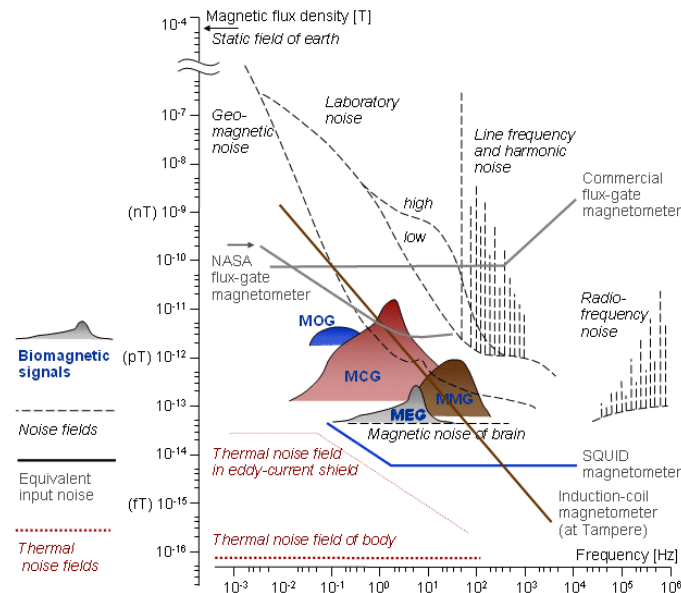
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Chapter 2: Principle description of Modulated Magnetic Signature of Blood

1 Introduction

Based on the known methods discussed in Chapter 1 for blood pulse acquisition, it can be concluded that there is a need for a novel method of blood pulse sensing that is low power, small and is able to acquire blood pulse independent of the electrical or optical contact with the skin. Such a method will support the development of pulse sensing devices that can be embedded in lifestyle devices such as massage chairs, watches and furniture for continuous acquisition of physiological vital signs.

In order to achieve blood pulse sensing that is independent of the electrical or optical contact with the skin, sensing of the bio-magnetic field generated from the blood circulatory system is one of the potentially solutions. However, such bio-magnetic fields are generally weak and will require magnetic sensors, such as SQUID, that are highly sensitive. In addition, environmental and earth magnetic field isolation are necessary. Therefore, there are challenges for such a system to achieve portability and low power (Figure 7).



Biomagnetic signals: MCG = magnetocardiogram, MMG = magnetomyogram, MEG = magnetoencephalogram, MOG = magneto-oculogram

Noise fields: static field of the Earth, geomagnetic fluctuations, laboratory noise, line frequency noise, radio frequency noise

Equivalent input noise: commercial flux-gate magnetometer, ring-core flux-gate (NASA), induction coil magnetometer, SQUID-magnetometer

Thermal noise fields: eddy current shield, the human body

Figure 7 Strength of magnetic signals produced by various sources [1]

In this chapter, the patented method [2] of blood pulse sensing using magnetic disturbance will be used to demonstrate blood pulse sensing that is independent of the electrical or optical contact with the skin. This method is based on the detection of magnetic disturbance when blood flows through a uniform magnetic field. An experimental setup will also be presented for data acquisition and analysis.

Data acquired in these experiments will be processed and the waveform will be analyzed in comparison with ECG and pulse plethysmography measurements acquired concurrently. In addition, the waveform acquired using the method of magnetic disturbance will also be compared with published waveforms such as MCG and analyzed relative to the activities of the heart. Finally, the conclusion section will summarize the work done and presents the feasibility of applying the patented method of blood pulse sensing to develop a small and low power system that is able to sense blood pulse independent of the electrical or optical contact with the skin.

2 Microelectronic overview on sensor

The magnetic sensor used in this dissertation to detect magnetic disturbance due to blood flow is based on the Giant-Magneto-Resistance (GMR) effect. As such, this section shall provide an overview on the materials used to construct the GMR sensors, the magnetic characteristics and its applications.

2.1 Introduction

In 1988, scientists discovered the “Giant-Magneto-Resistive” effect where a large change in electrical resistance occurs when thin, stacked layers of ferromagnetic and non-magnetic materials are exposed to a magnetic field. Since then, many companies have sought to develop practical applications for this intriguing technology. Examples of applications for the GMR effect are widely known in the hard-disk [3] and flash based memory industry [4], [5]. In addition, due to the high sensitivity of GMR sensor, applications of GMR sensor for bio-signal acquisition have also been reported [6],[7]. As such, based on the prior work done in the application of GMR sensor for biosignal acquisition and the merits described in Figure 8, the GMR based magnetic sensor was selected for this research.

Benefits:	GMR	HALL	AMR
Physical Size	Small	Small	Large
Signal Level	Large	Small	Medium
Sensitivity	High	Low	High
Temperature Stability	High	Low	Medium
Power Consumption	Low	Low	High
Cost	Low	Low	High

Figure 8 Comparison of GMR, Hall⁵ and AMR⁶ based magnetic sensor

2.2 Materials, structure and magnetic characteristics of GMR sensor

GMR is a quantum mechanical magnetoresistance effect observed in thin film structures composed of alternating ferromagnetic and nonmagnetic layers. GMR films have two or more magnetic layers separated by a nonmagnetic layer. Because of spin-dependent scattering of the conduction electrons, the resistance is maximum when the magnetic moments of the layers are antiparallel [8][9][10] and minimum when they are parallel. Since the discovery of GMR a large number of magnetic multilayer structures, which display the GMR effect, have been discovered. It was found that the magnitude of GMR varies considerably depending on the chemical constituents of the multilayer. Published values of high GMR are 220% in Fe/Cr multilayers [11] and 120% in Co/Cu multilayers[12]. Sizeable values of GMR were also obtained in multi-layers with Co/Ag – 22% at room temperature (RT) [13], Ni/Ag – 28% at 4.2K [14] and Ni/Cu – 9% at 4.2K [15].

The 3 commonly used structures in the construction of GMR sensors are unpinned sandwiches, antiferromagnetic pinned spin valves and antiferromagnetic multilayers [16]. Each of these structure has characteristic value usually associated with the GMR effect, which is the percent of change in resistance normalized by the saturated, or minimum, resistance.

Unpinned sandwich GMR materials consist of two soft magnetic layers of iron, nickel, and cobalt alloys separated by a layer of a nonmagnetic conductor (e.g., copper). With magnetic layers 4–6 nm (40–60 Å) thick separated by a conductor layer 3–5 nm thick, little magnetic coupling occurs

⁵ The Hall effect is the production of a potential difference (the Hall voltage) across an electrical conductor, transverse to an electric current in the conductor and a magnetic field perpendicular to the current and a transducer that is used to measure magnetic field using this phenomena is the Hall effect sensor.

⁶ Anisotropic Magnetoresistive (AMR) sensor is characterized by its simplicity, which consists of a thin permalloy layer and metal stripes (so-called barber poles) which cause a linearization of the sensor characteristics. The spontaneous magnetization lies in the easy axis direction which is fixed by shape anisotropy. A magnetic field along the heavy axis (perpendicularly to the easy axis) provides a rotation of the magnetization in the permalloy strip and thus a change of its resistance.

between the layers. Typically, sandwich materials have GMR values of 4%–9% and saturate with 2.4–5 kA/m (30–60 Oe) applied field [16].

Spin valves, or antiferromagnetically pinned spin valves, are somewhat similar to unpinned spin valves or sandwich materials. An additional layer of antiferromagnetic material is added at the top or the bottom. The antiferromagnetic material (e.g., FeMn or NiO) couples with the adjacent magnetic layer and pins it in a fixed direction. The other magnetic layer is free to rotate. If the external magnetic field is applied in a direction parallel to the magnetization of the pinned layer, there is a change in sheet resistance from its high level for one field direction to a low level for the opposite field direction (see Figure 9(a)). If the field is applied perpendicular to the pinned layer, the sheet resistance is minimum at zero field and increases for both positive and negative applied fields (see Figure 9(b)). The maximum change in sheet resistance in this configuration is only one half the total possible values. The free magnetic layer rotates from parallel to the pinned layer to perpendicular to it rather than from parallel to anti-parallel. GMR values based on this structure are typically 4%–20%, and the saturation fields are 0.8–6 kA/m (10–80 Oe) [16].

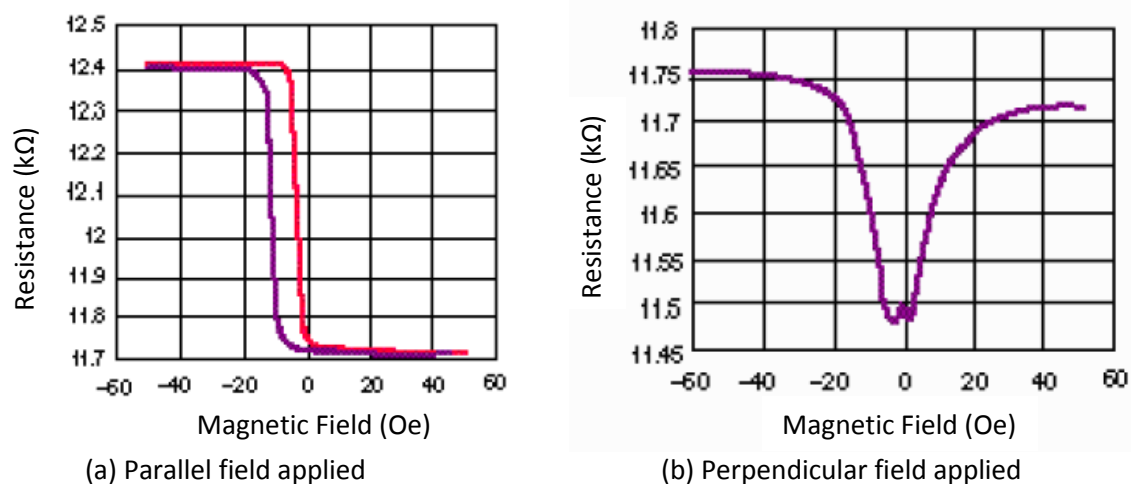


Figure 9 Plot of resistance vs applied magnetic field for an antiferromagnetically pinned spin valve GMR device with magnetic field applied parallel (a) and perpendicular (b) to the magnetization of the pinned layer [16]

Antiferromagnetic multilayers of GMR material consist of repetitions of alternating conducting magnetic layers and conducting nonmagnetic layers. Because multilayers have more interfaces than do sandwiches, the GMR effect is greater. The thickness of the nonmagnetic layers is less than that of the sandwich material (which is typically 1.5–2.0 nm), and the thickness is critical. The polarized conduction electrons cause anti-ferromagnetic coupling between the magnetic layers only when the spacers are a certain thickness. In the absence of an external magnetic field, each magnetic layer has its magnetic moment anti-parallel to the moments of the magnetic layers on each side—exactly the condition needed for maximum spin-dependent scattering. A large external field can overcome the

coupling that causes this alignment and can align the moments so that all the layers are parallel--the low resistance state. If the conducting layer is not the proper thickness, the same coupling mechanism can cause ferromagnetic coupling between the magnetic layers resulting in no GMR effect.

A plot of resistance vs. applied field for a multilayer GMR material is shown in Figure 10. Note the higher GMR value, typically 12%–16%, and the much higher external field required to saturate the effect, typically 20 kA/m (250 Oe). Most multilayer GMR materials have better linearity and lower hysteresis than sandwich GMR material [16].

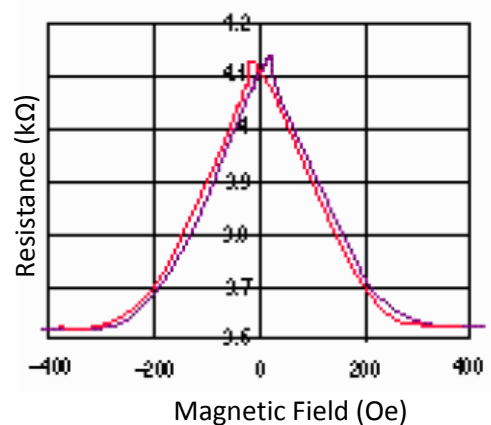


Figure 10 Plot of resistance vs. applied field for a 2 mm wide strip of anti-ferromagnetic coupled multilayer GMR material [16]

2.3 Selection of GMR sensor for research

Based on the characteristics of the various GMR structure presented, the spin valves structure was selected based on its higher sensitivity and the availability of a commercial solution to facilitate the research on the application of GMR sensor for magnetic blood pulse sensing.

3 Basic principles on Modulated Magnetic Signature of Blood (MMSB)

In this dissertation, the term used to describe the phenomena of magnetic disturbance created by blood pulse flowing through an applied magnetic field is termed Modulated Magnetic Signature of Blood (MMSB).

MMSB can be illustrated in Figure 11 where a uniform magnetic field is applied on the skin, in a non-invasive manner, within close proximity of a major blood vessel and a magnetic sensor. The uniform magnetic field can be generated from magnet sources such as a small permanent magnet or electromagnet with magnetic field strength of 0.1 - 0.2 Tesla. The applied magnetic field will create a uniform magnetic field encompassing the Giant-Magneto-Resistance (GMR) based magnetic sensor,

skin, fabric and blood vessel shown in Figure 11. Such a magnetic field, when undisturbed, will produce a fixed DC output voltage from the magnetic sensor based on the sensor characteristics as illustrated in Figure 12. In addition, the technique of applying the magnetic biasing field also creates a biasing field to bias the sensor away from the zero field area, where hysteresis is more pronounced.

The application of magnetic field within close proximity of the GMR sensor also describes the phenomena of magnetic biasing, which will operate the GMR sensor in the linear region. Therefore, the magnetically biased sensor will be able to detect and amplify the weak magnetic disturbance or MMSB created by the pulsatile flow of blood through the uniform magnetic field.

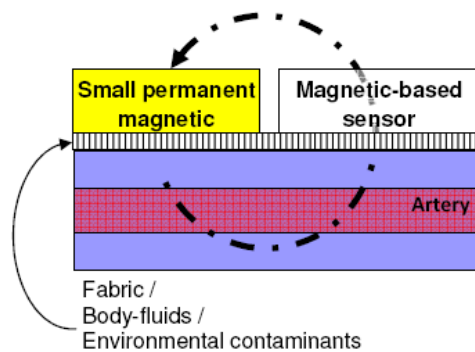


Figure 11 Cross-sectional view illustrating the two-dimensional concept for the experimental setup to acquire MMSB

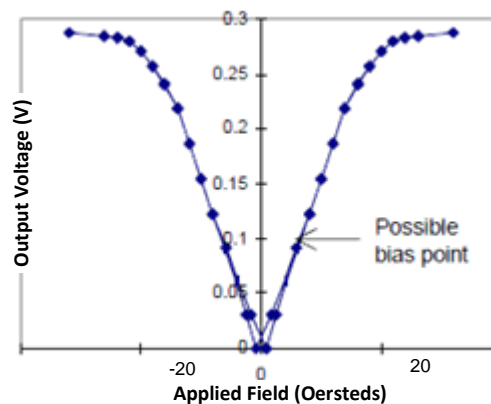


Figure 12 Typical characteristics of GMR sensor (in Oersteds⁷) used in this experiment

3.1 Applications of GMR sensor for magnetic blood sensing

The GMR sensor used in this research work is based on the GMR sensor provided by the company NVE Corporation. Based on the datasheet provided for the GMR sensor [17], it was found that the sensor is not sensitive enough to acquire the bio-magnetic signal from the human blood without amplifying the bio-magnetic signal. As such, the concept of biasing magnetic field was applied to achieve amplification effects. Biasing magnetic fields will provide either a magnetic field to sense (where one may not be present) or create a pseudo field for the sensor. Such a biasing magnetic

⁷ 1 Tesla = 10,000 Oersteds

field typically describes the application of a magnetic field through the sensor package without influencing the sensor operations. An example of a commercially available application using magnetic biasing technique is to detect the ferrous material of currencies so as to identify and differentiate the notes automatically [18], [19].

To estimate the distance between the permanent magnetic and the GMR sensor to achieve the desired effect of magnetic biasing, the magnetic field at the sensor location will be estimated. In order to achieve this estimate, the permanent magnet is assumed to be a point source and the magnetic field, B along its axis, at distance d away from the magnet can be calculated using Equation 6 (reference [20]).

Equation 6
$$B = \frac{\mu_0 \mu}{4\pi d^3} (a_R 2\cos\theta + a_\theta \sin\theta)$$

where μ_0 = permeability constant ($4\pi \times 10^{-7}$ T m/A⁸)

d = distance from the center of the magnet in meters

a_R and a_θ = spherical unit vectors from centre of magnet (Figure 13)

μ = magnetic moment of a magnet (A.m²).

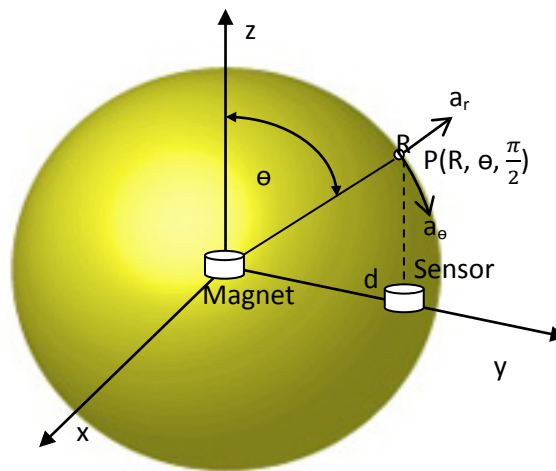


Figure 13 Illustration of magnetic field vector calculations based on [21]

Applying Equation 6 with $\theta = 90^\circ$, the magnetic field B can be simplified to Equation 7.

Equation 7
$$B = \frac{\mu_0 \mu}{4\pi d^3} (a_\theta)$$

Using Equation 7 and applying the physical data where $d = 0.02\text{m}$, magnetic moment of the magnet $\approx 0.05 \text{ Am}^2$, the magnetic field strength at the sensor d away from the magnet can be calculated as 6.25 Oe , where 1 Oe is approximated as 1 Gauss (or $1 \times 10^{-4} \text{ T}$). This will produce the effect of

⁸ Where A is a vector magnetic potential with units Wb/m

magnetic biasing where the sensor is always operating in the linear region of its characteristic curve as shown in Figure 12.

3.2 Experimental setup to demonstrate MMSB using iron oxide fluidic flow

To demonstrate the existence of a magnetic disturbance when blood flows through a constant magnetic field, an experimental setup will be described in this section to verify the MMSB phenomenon using iron (II) oxide (Fe_2O_3) fluid as shown in Figure 14.

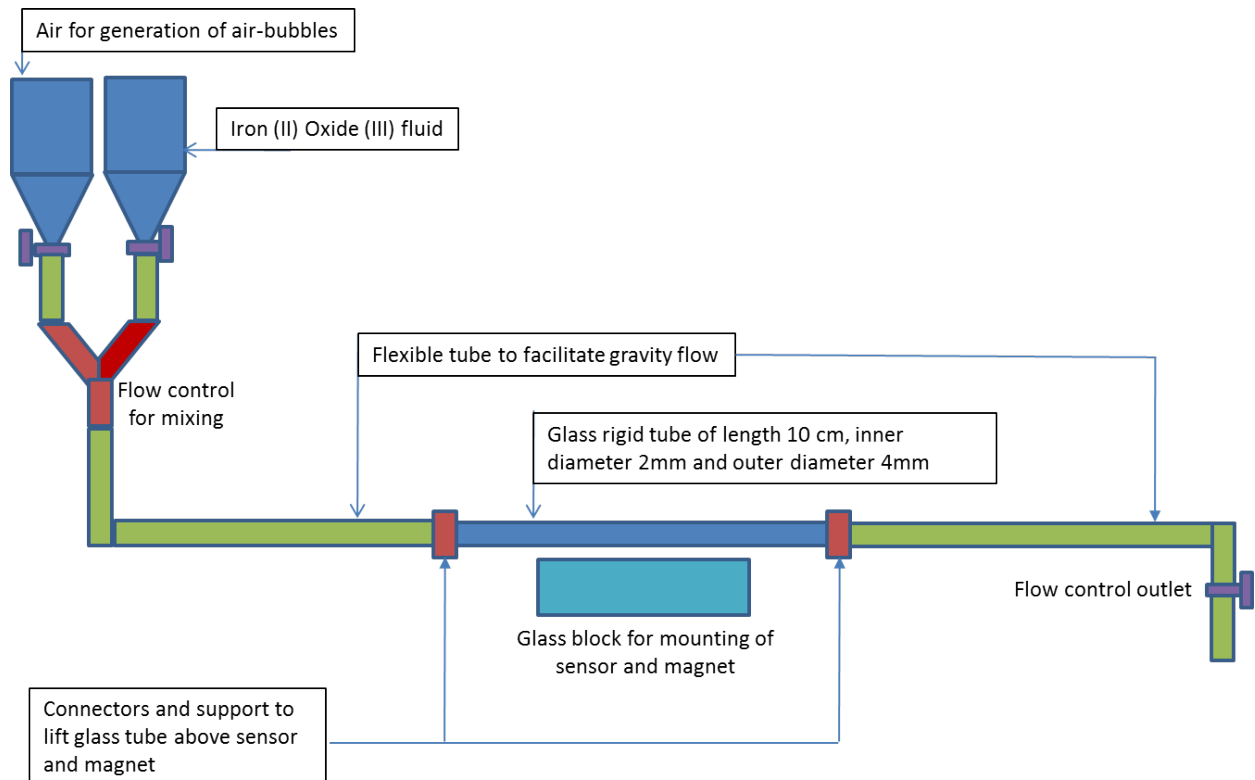


Figure 14 Iron (II) oxide (Fe_2O_3) fluidic flow experiment

The experimental setup on Figure 14 is designed to make use of gravitational force to allow the Fe_2O_3 fluid to flow through the rigid glass tube, which is placed at 2-3mm above the sensor and magnet. However, continuous flow of Fe_2O_3 fluid lacks the ability to demonstrate the effect of MMSB, which is primarily observed when pulsatile blood flows through a constant magnetic field. Therefore, an extra funnel is used in Figure 14 to support inlet of air into the fluidic flow so that pockets of air-bubbles can be infused between the Fe_2O_3 fluids as it flows through the system. This is illustrated in Figure 15, where the effect of magnetic disturbance is tested through flow of Fe_2O_3 fluids that is separated by pockets of air-bubbles.

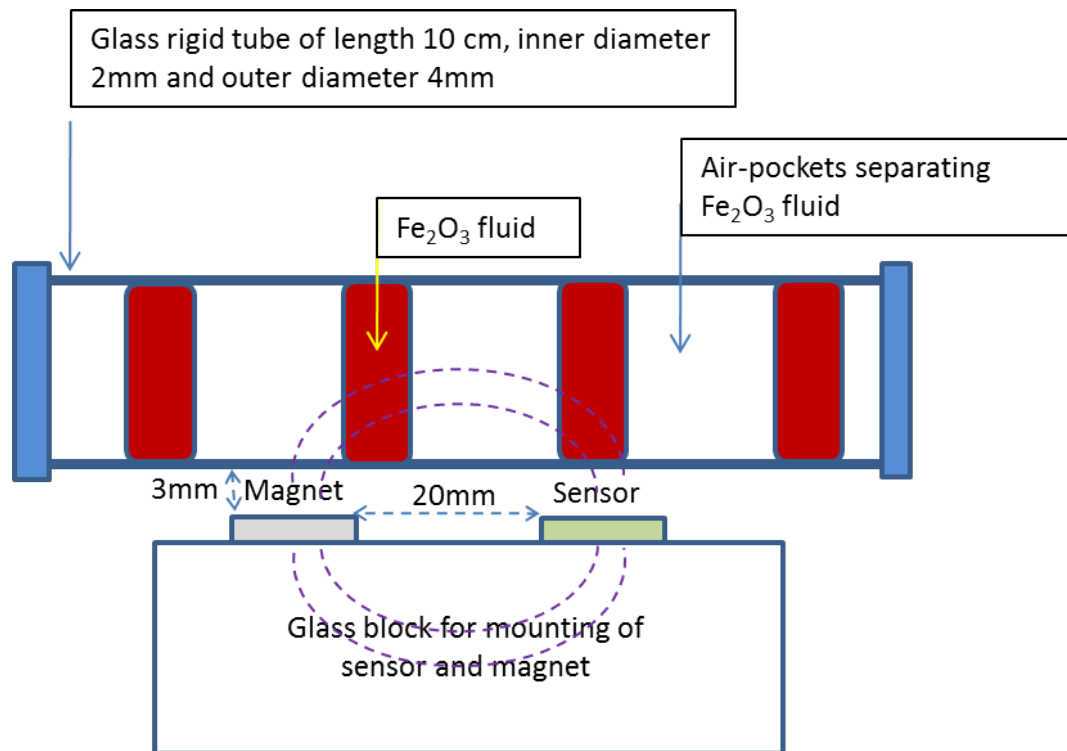


Figure 15 Illustration on flow of Fe_2O_3 fluids separated by pockets of air-bubbles

Using this setup shown in Figure 14, the inlet flow control of Fe_2O_3 fluids is controlled together with the flow outlet control valve such that the physical generation of Fe_2O_3 fluidic flow separated by pockets of air-bubbles is formed as shown in Figure 15. As such, a mechanical pump is not used in this experiment to simulate the pulsatile effect of blood flow.

In addition, the rate of Fe_2O_3 fluid flow is controlled using manual adjustment on inlet and outlet valves. Therefore, the rate of Fe_2O_3 fluid flow is not the same as that of the human blood flow. However, the experiment is sufficient to be used as a proof-of-concept for the proposal of magnetic disturbance in MMSB.

The final laboratory setup for this experiment is shown in Figure 16, where the sensor output is amplified by a differential amplifier with gain of 100. The amplified signal is connected to the National Instrument data acquisition module NI USB-6008, which samples the signal using a 12 bit Analog to Digital Conversion (ADC) with sampling rate of 1 kHz. As shown in Figure 16, all measured data are concurrently displayed and stored on the laptop.

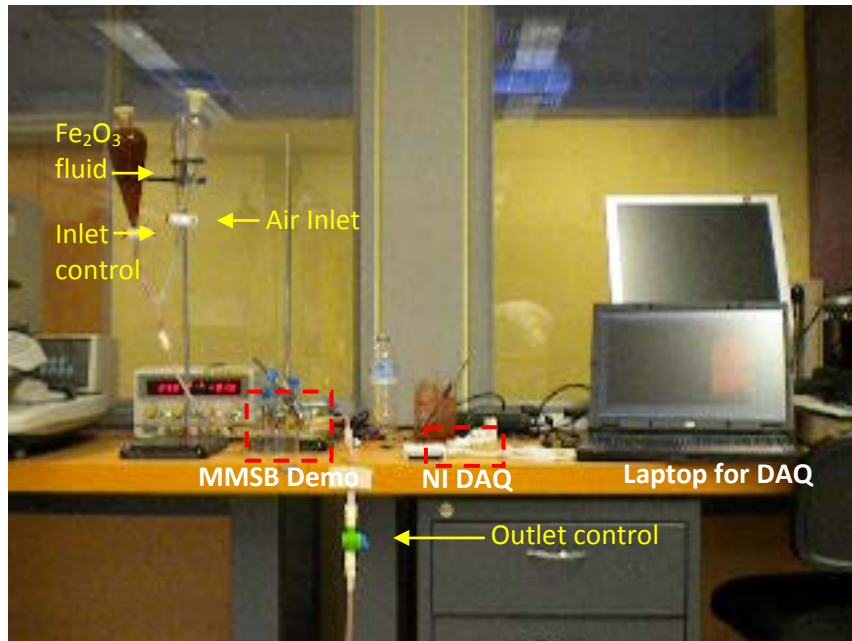


Figure 16 Experimental setup used in laboratory

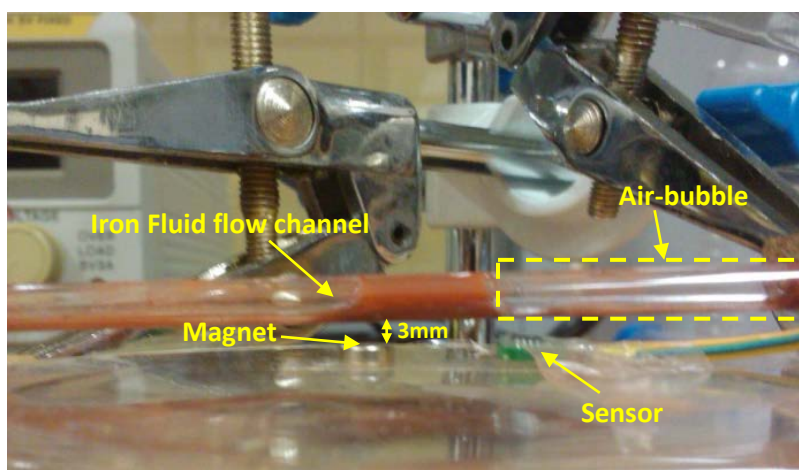


Figure 17 Position of magnet and sensor with respect iron fluidic flow channel

3.3 Measurement results for Fe_2O_3 fluidic flow experiment

The measurement result for Fe_2O_3 fluidic flow is obtained based on experiments conducted using Fe_2O_3 concentration as a percentage of haemoglobin in blood (taken as 16 g/dL for calculation purpose), with reference to published results (<http://www.bloodbook.com/ranges.html>), where range of haemoglobin in male is estimated as 13 - 18 g/dL and in female it is estimated as 12 - 16 g/dL. The results of this experiment are tabulated in Table 3 with samples of waveform obtained shown in Figure 18.

S/N	Fe ₂ O ₃ concentration (g/dL)	% of Fe ₂ O ₃ concentration used with reference to 16g/dL of Haemoglobin	Measured peak-to-peak voltage (mV)
1	16	100.00%	58
2	12	75.00%	51
3	8	50.00%	43
4	3.61	22.56%	32
5	1.33	8.31%	18
6	0.83	5.19%	12
7	0.41	2.56%	9

Table 3 Measurement result for Fe₂O₃ fluidic flow in constant magnetic field

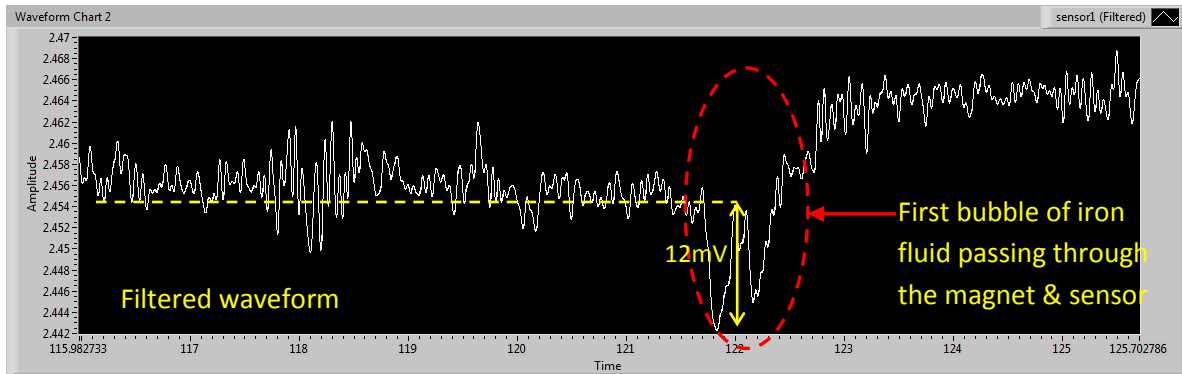


Figure 18 Waveform obtained when Fe₂O₃ fluid flow pass the sensor with air-pockets

One of the challenges in this experiment is the precipitation of iron oxide above the magnet as shown in Figure 19. Therefore, the flow rate in each experiment has to be controlled manually so that the precipitation of iron oxide does not affect the measurement results. In addition, upon the completion of each experiment, the setup has to be flush with continuous flow of water without the magnet in place.

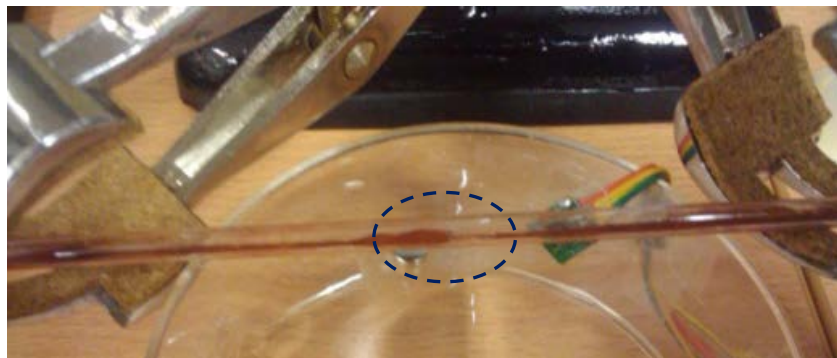


Figure 19 Precipitation of Fe₂O₃ above magnet

The experiments were successfully carried out using the manual control of the inlet and outlet valves to achieve iron fluid flow with air bubbles as shown in Figure 20. A typical sample of the signal acquired is shown in Figure 21, where magnetic disturbance can be observed during iron fluid flow with air-bubble.

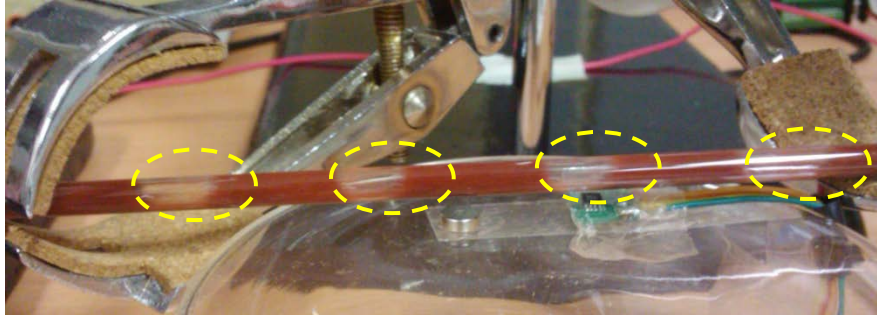


Figure 20 Flow of iron fluid with air-bubbles

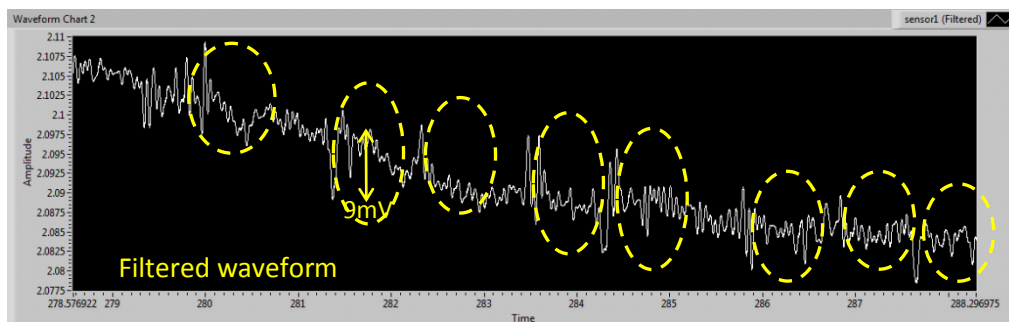


Figure 21 Signal acquired during iron fluid flow as shown in Figure 20

With the successful completion of these experiments, the measured results demonstrate the presence of magnetic disturbance when iron oxide fluid flows through a constant magnet field. As such, the basis of MMSB was demonstrated successfully through the use of a laboratory setup using Fe_2O_3 fluid. The following sections will describe experiments conducted on humans to measure the magnetic disturbance when blood flows through a constant magnet field.

3.4 Acquisition of magnetic disturbance of blood

With the successful demonstration of the existence of a magnetic disturbance due to the flow of a ferrous fluid, this section will outline the experiment to acquire the magnetic disturbance created due to the pulsatile blood flow of blood. The magnetic disturbance will be amplified by the magnetically biased GMR sensor allowing MMSB to be measured at room temperature.

Based on the principle of applying an external magnetic field to penetrate a thin layer of material (example: human skin) that is non-magnetic, MMSB can be measured non-invasively. In addition, this layer of non-magnetic material may also include fabric, bodily fluids and environmental

contaminants (Figure 11), allowing MMSB to have unique advantage over existing heart rate measurement systems.

Using the blood circulatory system for human as shown in Figure 22, it can be observed that there are many locations where MMSB signal acquisition is possible. These locations are characterized with major blood vessels (particularly blood vessels with oxygenated blood), easily accessible (with minimum hair) and have minimum human tissues such as fats and muscles. Examples of such locations where MMSB can be easily acquired are the wrist, heel, temple and neck as shown in Figure 22.

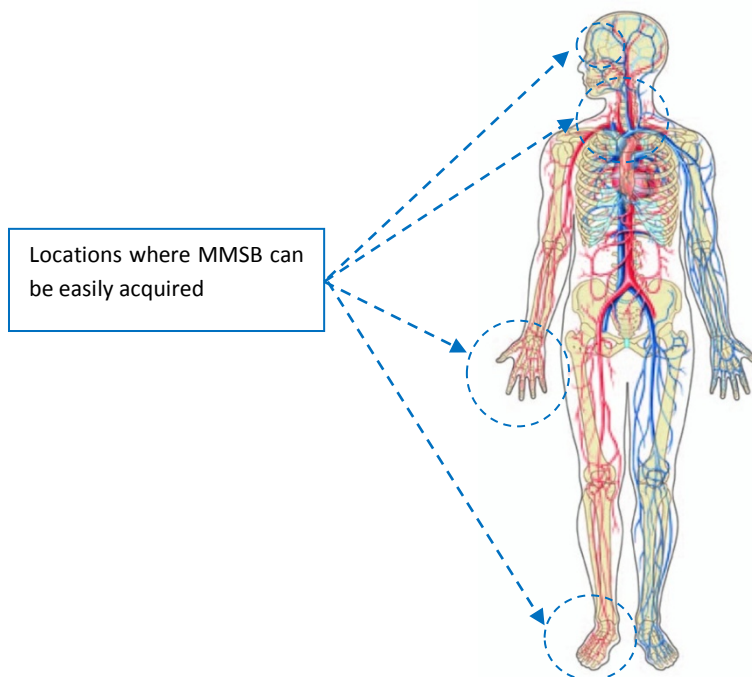


Figure 22 Illustrations of example locations where MMSB can be acquired easily on the Human Blood Circulatory system

Experiments were setup and carried out on the locations labeled in Figure 22 to verify the hypothesis of MMSB. Based on measured results, MMSB data was successfully collected on all these locations. However, during the experiment, it was found that measurement on the temple is not feasible for trials on human subjects. This is because the GMR sensor and magnet needs to be mounted vertically on the subject's face (as shown in Figure 23) using adhesive tapes and each trial will result in the hair of the subject being removed.

During the experiment to collect MMSB data on the heel, it was concluded that the heel is not a comfortable location for extensive MMSB measurement. This is because the heel is not easily assessable without the subject feeling uncomfortable (as shown in Figure 24).

Finally, experiment work to collect MMSB data on the wrist was found to be most appropriate. This is because the wrist has minimum hair and most subjects are able to rest their arm comfortably during the experiment. As such, the following sections will describe experimental setup and results based on MMSB acquired on the wrist (as shown in Figure 25). Using experimental results obtained from measurements on the wrist, the optimal distance between the sensor and magnet is found to be applicable on other locations such as the heel, neck and temple.

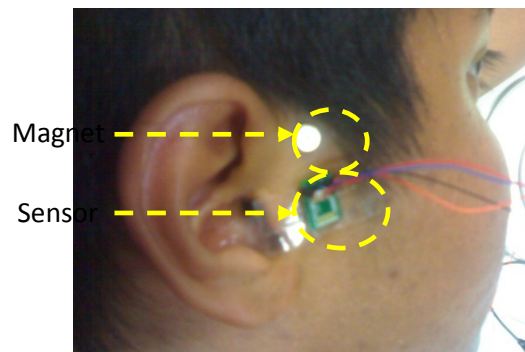


Figure 23 Illustration of sensor and magnet mounting near temple for MMSB signal acquisition

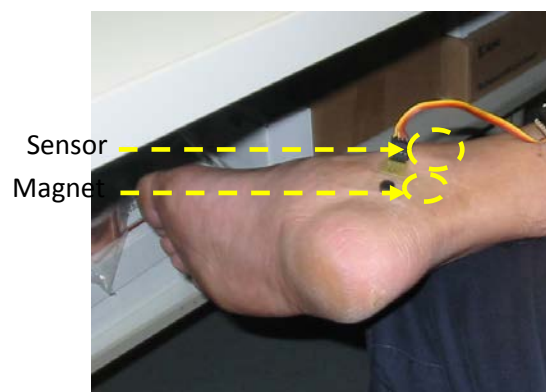


Figure 24 Illustration of sensor and magnet mounting on the heel for MMSB signal acquisition

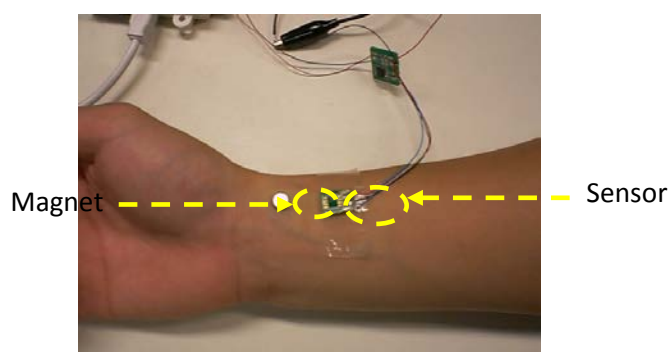


Figure 25 Illustration of sensor and magnet mounting on the wrist for MMSB signal acquisition

4 Experimental setup for MMSB signal acquisition and analysis

The experimental setup for MMSB signal acquisition is based on measurements on the wrist where further work in Chapter 6 will demonstrate the applications of the optimal distance between sensor and magnet, found in this experiment, to acquire MMSB on the heel (Figure 26(b)).

Based on the discussions presented in Section 3.2 of Chapter 2, the wrist is chosen for this experiment due to its easy access with minimum human tissues and hair. In addition, the major artery on the wrist can be easily found, allowing experiments to be conducted efficiently.

The experimental setup is illustrated in Figure 26(a), where the relative position between the magnet (1), sensor (2) and the artery (major blood vessel) on the wrist is varied to achieve maximal output voltage from the sensor. With reference to the cross-sectional view illustrated in Figure 11, the distance between the magnet and the sensor is measured for each measurement and the result is correlated to the measured amplitude of the MMSB signal. The amplitude of the MMSB signal is obtained using the system block diagram as illustrated in Figure 27 and the circuit used is shown in Figure 28. The differential signal from the sensor is amplified using a differential to single ended amplifier with gain of 100 times. The amplified signal is connected to an oscilloscope and the National Instrument data acquisition module NI USB-6008 that is integrated with 12 bit Analog to Digital Conversion (ADC) with sampling rate of up to 10 kS/s. In this experiment, the signal is sampled at 1 kHz, which is 10 times higher than the typical ECG sampling rate. The digitized signal is sent to a computer for pulse rate measurements and the results are compared with the pulse rate measured by the pulse oximeter, which is connected to the finger of the human subject.

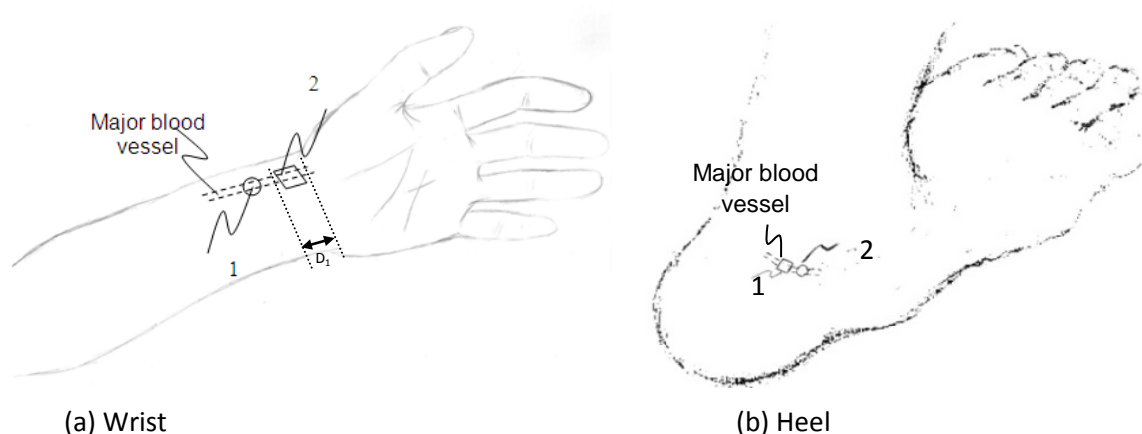


Figure 26 Illustrations of the relative position of the magnet(1), sensor(2) and a major blood vessel

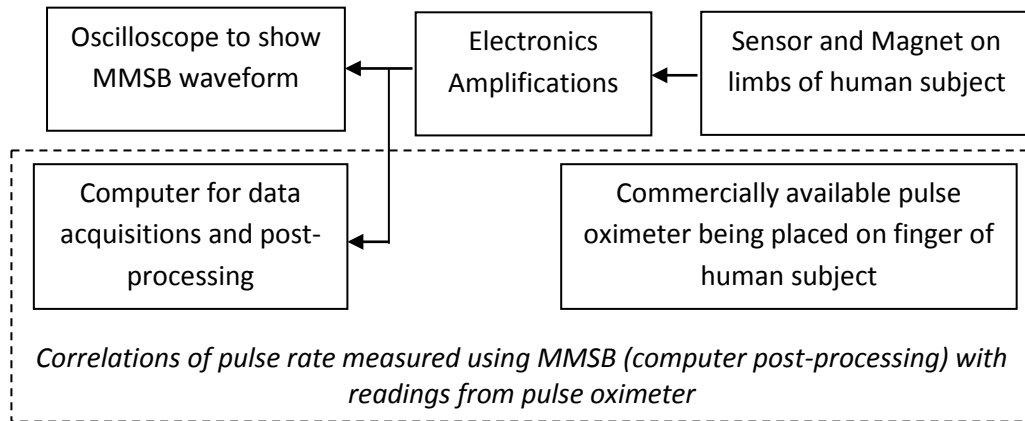


Figure 27 System Block Diagram used for MMSB signal acquisition

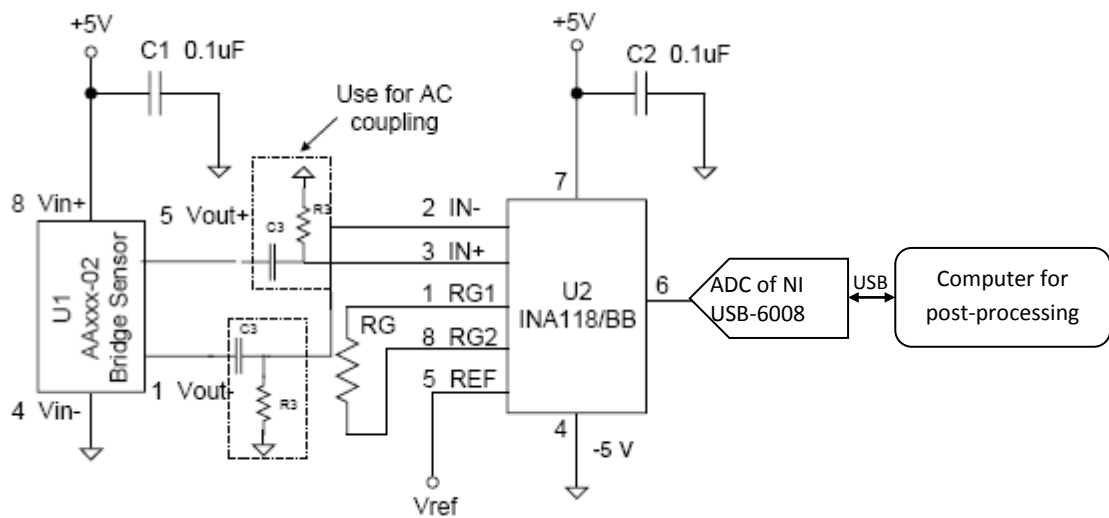


Figure 28 Circuit used for MMSB signal conditioning

This experiment is setup to operate at room temperature using a button-sized permanent magnet such as the Sola 1000 [22] or the Assemtech - M1219-4 [23] or any equivalent magnet with approximately 1000 Gauss and a diameter of 6mm. The magnetic field strength of the permanent magnet has to be controlled to be within the safety range for human beings as described by The International Commission on Non-Ionizing Radiation Protection (ICNIRP) Guidelines [24].

Based on the ICNIRP guidelines, the human continuous exposure to occupational static magnetic fields should be limited to be less than 0.2 Tesla. For short-term whole-body exposure, the recommended magnetic field strength must be less than 2 Tesla. Finally, for short-term arms and legs exposure, the recommendations are to limit the magnetic field to less than 5 Tesla. However, for persons with pacemakers and other implanted devices, ICNIRP recommends the magnetic field exposure to be limited to a much lower level at 0.5 milliTesla (or 5 Gauss). This level is not related to biological effects but rather to possible effects on electrical or electronic devices (particularly reed-relays) or metal prosthesis.

Therefore, the magnetic field strength selected for this experiment is less than 0.2 Tesla (i.e. magnet \approx 1000 Gauss). This will facilitate both long and short-term measurements allowing continuous measurements of blood pulse for human subject.

The magnet selected is placed on the major arteries on the wrist with a commercially available Giant Magneto-Resistor (GMR) magnetic sensor [17] placed in close proximity on the human skin. The measurements will depend on several aspects: the distance between the sensor and the magnet (approximately 20 mm) providing a uniform magnetic field that ensures proper biasing, the magnet strength that does not saturate the sensor, and the appropriate penetration of the magnetic field into the skin tissues.

Using the experimental setup described above, measurements are conducted on more than 20 human subjects where MMSB waveforms were measured from all subjects. A typical MMSB waveform is shown in Figure 29 and it can be observed that the waveform is highly periodic. The periodicity of the waveform is found to be similar to the heart rate measured by the electrocardiogram or pulse plethysmography.

The characteristics of the waveform and its pulse width will be further analysed in later sections where comparison to published results will be presented with correlations to the activities of the human heart.

Having successfully setup the experiment to acquire MMSB on human subjects, the following sections will include experimental setups used to demonstrate the ability of MMSB to acquire blood pulse through fabric, blood and underwater. The successful demonstrations of these measurements will allow MMSB to be deployable for various applications including some of those described in Chapter 6.

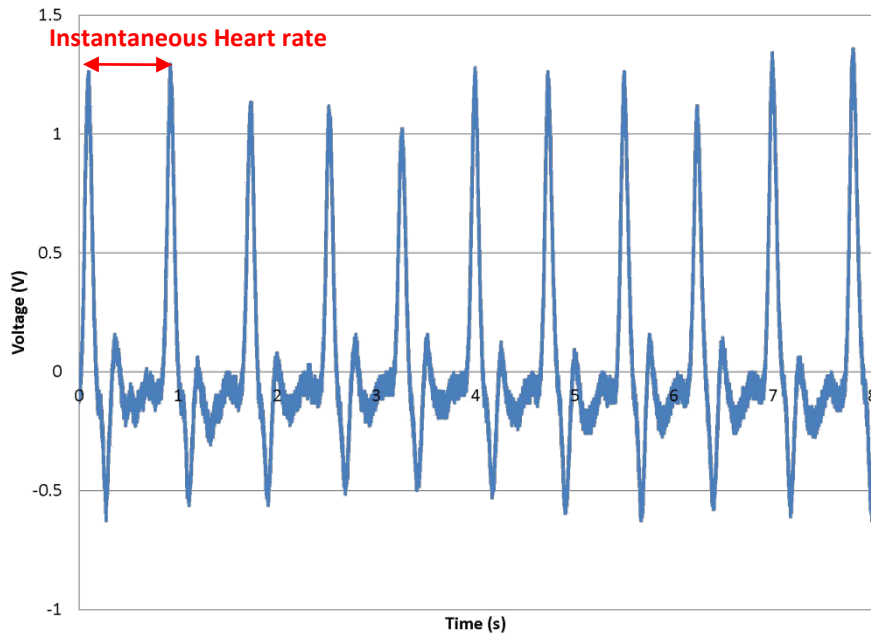


Figure 29 Waveform captured from the amplifier output using oscilloscope

4.1 Experimental setup to measure blood pulse through fabric

The objective of this experiment is to demonstrate the ability of MMSB in acquiring heart rate of measurement subject who have light bandage.

Using the basic principle shown in Figure 11, the experimental setup is illustrated in Figure 30 where the subject is lightly bandaged on the wrist (approximately 3mm thickness) with the GMR sensor and magnet mounted on top of the bandages using a strip-on configuration. Data is sampled at 10Hz with 10 bit resolutions and sent through the XBee® ZB ZigBee® PRO Radio Modems [26]. This modem is based on the ZigBee protocol and has an integrated Analog-to-Digital convertor (ADC) to sample the signal. However, the maximum sampling rate supported is 1 kHz and the memory available for storage before transmission is 46 samples. As such, in order to optimize the RF bandwidth, the ADC output is stored before transmission. Therefore, for this experiment, a sampling rate of 10 Hz is chosen and the schematic used for the transmitter is shown in Figure 31.

Using the ZigBee based wireless transmission system, data is received from the transmitter, processed and displayed on the computer using the block diagram shown in Figure 32.

Based on the data acquired using the MMSB phenomena and concurrent acquisition of pulse rate using the pulse plethysmography, the pulse rate of the subject is found to be similar for both methods of measurements. Therefore, it can be concluded that MMSB has been successfully demonstrated to acquire blood pulse through opaque objects (i.e. bandage) without the need of direct electrical or optical contact with the human skin.

Waveform of blood pulse at sampling rate of 10Hz

Wireless receiver

Sensing of blood pulse through light bandages with wireless data transmission

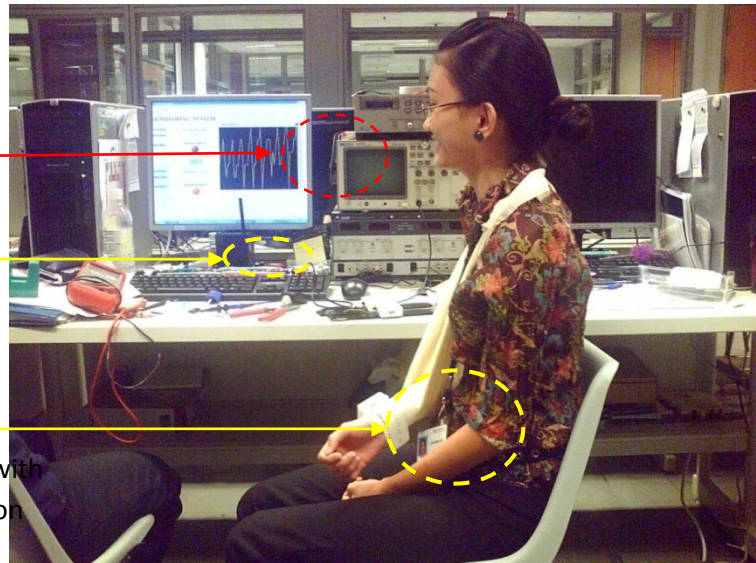


Figure 30 Illustration of blood pulse acquisition through light bandages

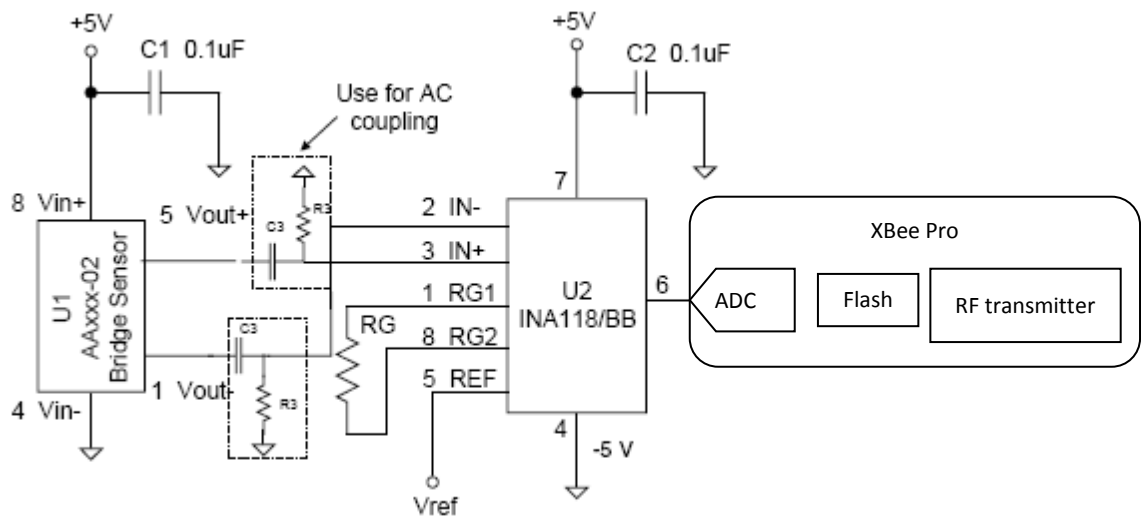


Figure 31 Schematic of MMSB signal acquisition, conditioning, sampling and transmission circuit

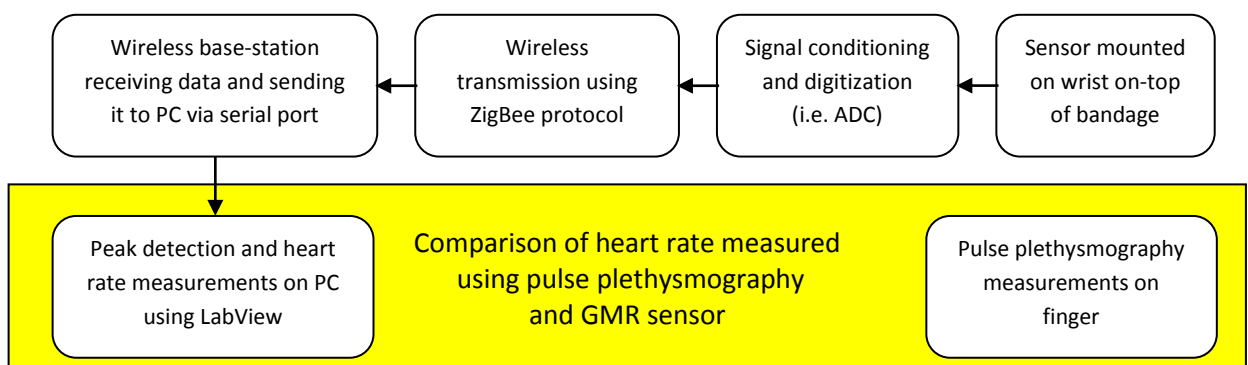


Figure 32 Block diagram of blood pulse signal acquisition and transmission back to base-station

4.2 Experimental setup to measure blood pulse through blood

The objective of this experiment is to demonstrate the ability of MMSB in acquiring heart rate of the subject who are injured and are bleeding. Heart rate for the injured person is an important physiological vital signs allowing the risk for any rescue operations to be assessed objectively.

The experimental setup is shown in Figure 33 where Figure 33(a) illustrates the procedure where blood is first smeared on the wrist of the subject. This is accompanied by Figure 33(b), where the sensor and magnet are mounted onto the blood stains. The oscilloscope in Figure 33(b) shows the successful acquisition of MMSB waveform from the subject. Therefore, it can be concluded that the MMSB phenomena is able to detect the heart rate through blood stain.

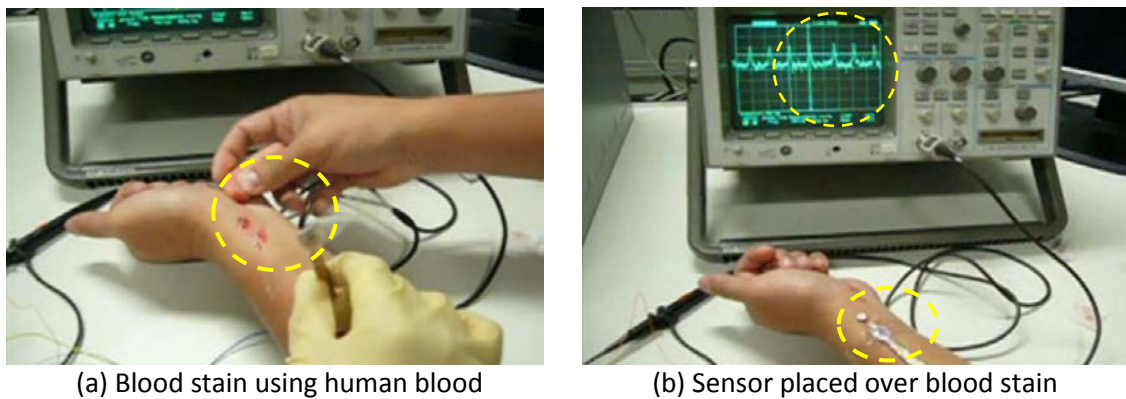


Figure 33 Illustration of blood pulse acquisition through blood stains

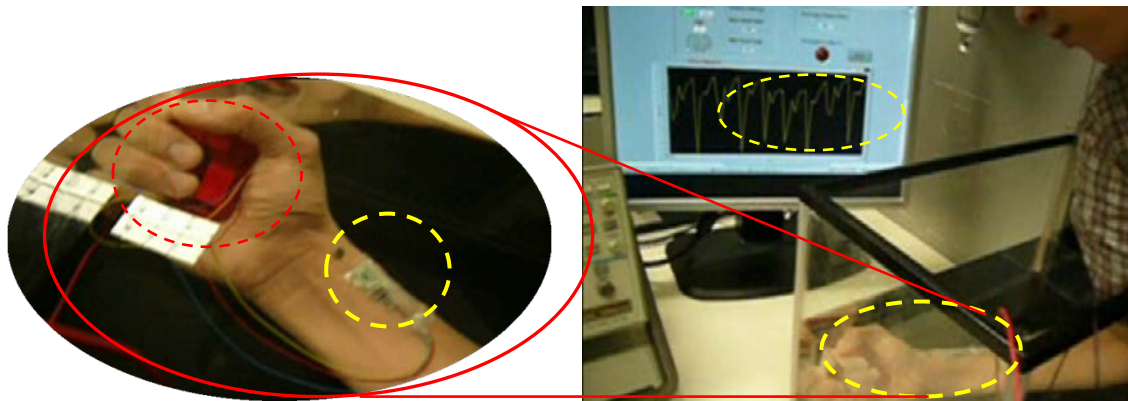
4.3 Experimental setup to measure blood pulse underwater

The objective of this experiment is to demonstrate the ability of MMSB in acquiring heart rate of subjects who are underwater or swimming. Heart rate for the divers or competitive swimmers is an important physiological vital sign allowing the risk for divers to be assessed continuously or the assessment of the athletic performance for swimmers.

The experimental setup is shown in Figure 34 where the subject's arm is immersed in a tank of water with the GMR sensor and magnet on the wrist (Figure 34(a)). Data is sampled at 10Hz with 10 bit resolutions and sent through a wireless transmission system using similar configurations as shown in Figure 31. Data collection, analysis and display are completed the computer with the results shown in Figure 34(b).

Based on the data acquired using the MMSB phenomena and pulse plethysmography, the heart rate of the subject was found to be similar. Therefore, this experiment concludes the successful acquisition of using the MMSB phenomena to measure the heart rate of subjects who are in the rain

or under-water and subjects who are perspiring. However, as this experiment is conducted with the subject in a stationary position, applications in such scenarios will require the subject to be at stationary for a momentary period so as to minimize motion artifacts in the pulse rate measured.



(a) Sensor in water with wireless transmitter (b) Blood pulse measurement in water

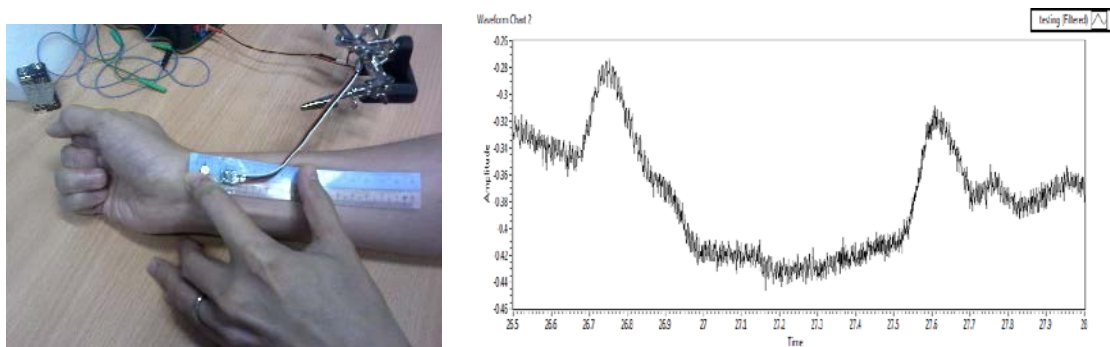
Figure 34 Illustration of blood pulse acquisition in water

4.4 Experimental setup to measure blood pulse through a plastic ruler

The objective of this experiment is to demonstrate the ability of MMSB in acquiring heart rate of subjects through a soft and flexible solid material such as plastic ruler.

The experimental setup is shown in Figure 35(a), where the placement of a plastic ruler between the sensor and the wrist of the human subject is illustrated. Using the signal conditioning and data acquisition schematic shown in Figure 31, the sensor output is monitored. The output of the data acquisition module is shown in Figure 35(b), which demonstrates the successfully acquisition of MMSB pulse data through a soft and flexible solid material such as plastic ruler.

Therefore, from this experiment it can be concluded that the MMSB phenomena can be detected through a thin layer of soft and flexible solid material such as a plastic ruler.



(a) Sensor and magnet placed on plastic ruler (b) Typical signal obtained with digital filtering

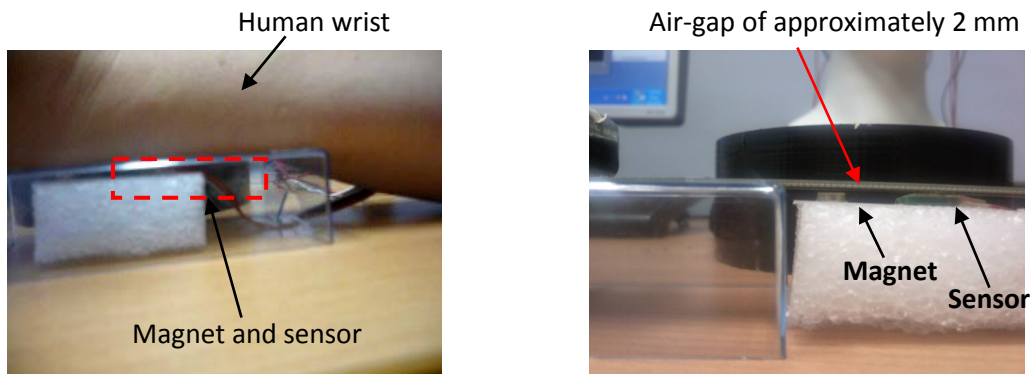
Figure 35 Blood pulse acquisition through plastic ruler placed in between sensor and wrist

4.5 Experimental setup to measure blood pulse through air-gap

The objective of this experiment is to demonstrate the ability of MMSB in acquiring heart rate of subjects through a small air-gap between the wrist and the sensor.

The experimental setup is shown in Figure 36(a), where the wrist of the human subject is rested on a plastic structure to ensure that the 2 mm air-gap shown Figure 36(b) is not bridged. Using the signal conditioning and data acquisition schematic shown in Figure 31, the sensor output is monitored. The output of the data acquisition module is shown in Figure 37, which demonstrates the successfully acquisition of MMSB pulse data through an air-gap as discussed in this section.

Therefore, from this experiment it can be concluded that the MMSB phenomena can be detected through an air-gap as described in Figure 36.



(a) Wrist placed on plastic structure to maintain air-gap (b) Sensor and wrist separated by air gap

Figure 36 Illustration of blood pulse acquisition setup through air-gap

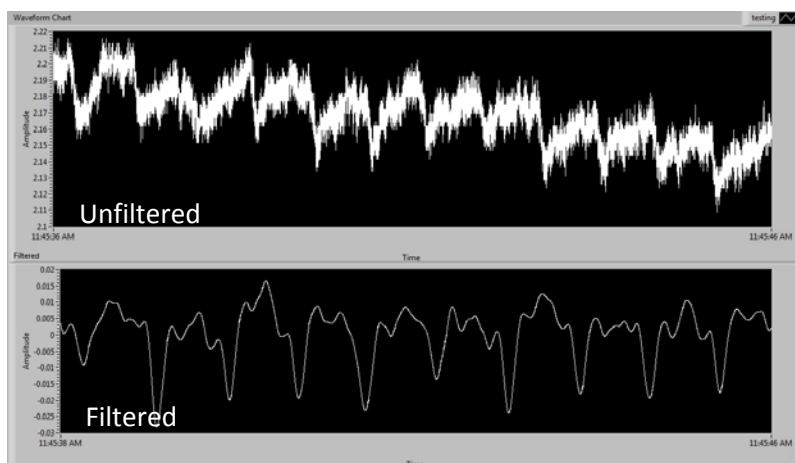


Figure 37 Sample MMSB waveform obtained using setup shown in Figure 36

5 Comparison of MMSB with ECG and pulse plethysmography waveform

The experimental setup for MMSB signal acquisition on the wrist is simultaneously acquired with the ECG and the pulse plethysmography using the USBamplifier from Guger Technologies (g.tec). The USBamplifier will digitize and display the waveform using MATLAB Simulink. The block diagram of the MATLAB Simulink setup used in this experiment is shown in Figure 38 with ECG, pulse plethysmography and MMSB signals being sampled and displayed. The measured waveforms are shown in Figure 40 using similar sampling rate and filter settings.

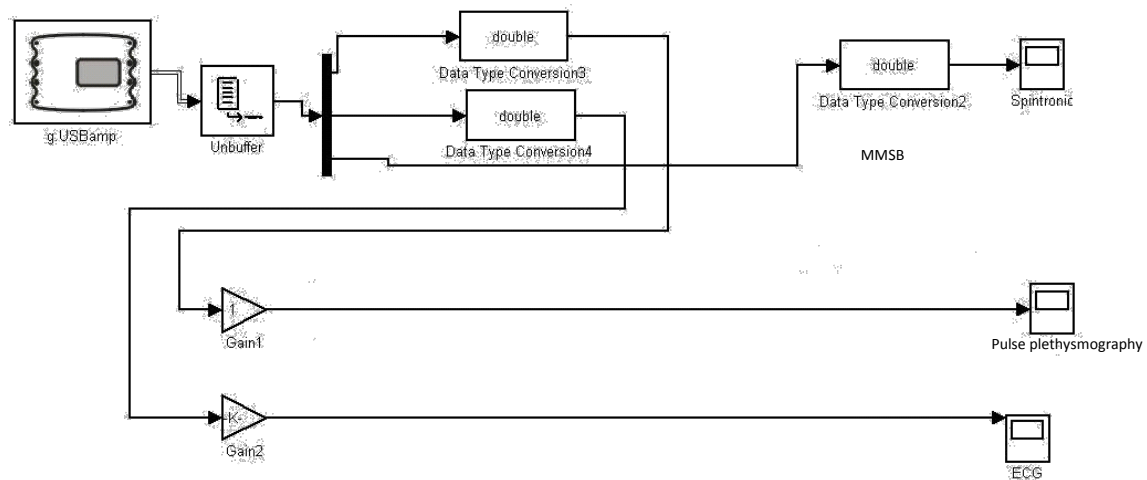


Figure 38 Block diagram of the MATLAB Simulink setup for MMSB signal acquisition on the wrist with ECG and pulse plethysmography using the USBamplifier from Guger Technologies

Based on the data acquired in Figure 40, it can be observed that the signal acquired from all the three methods produce similar periodicity with distinct waveform characteristics. The periodicity of the waveform indicates the heart rate of the subject while the characteristics of the waveform acquired are dependent on the method of signal acquisition.

The ECG is the golden method used for the medical industry for heart rate measurements. This is because ECG is a direct measurement of the electrical activity of the heart and is based on electrical signals acquired from the triangulation of electrodes placed around the heart. Therefore, the similarity in periodicity of the waveform acquired using the GMR sensor and ECG electrodes conclude the reliability and accuracy of the MMSB phenomena to measure heart rate.

In addition to the periodicity, it can also be observed that there is a phase difference between the waveform acquired using the GMR sensor and ECG electrodes. This is due to the difference in the method of measurements where the ECG measures the electrical activity of the heart (electrical activity of the heart before it pumps) while the GMR sensor measures the magnetic disturbance of

the blood flow (mechanical activity of the heart when it pumps). This delay is dependent on factors such as the heart functionality and heart rate, compliance of the arteries and the distance of the probe from the heart. It has been reported [27] that the delay between ECG and pulse plethysmography is approximately 100ms for healthy adults and this is applicable for ECG and MMSB waveform as pulse plethysmography output is found to be in-phase with MMSB waveform.

The pulse plethysmography is chosen for comparison in this experiment as it is a commonly available lifestyle product. As such, it is important that the waveform acquired by the GMR sensor has to be capable of producing similar periodicity and phase difference as compared to the pulse plethysmography. In addition, both methods have a common similarity in measuring the result of blood disturbance to an applied magnetic and optical field respectively. This is because the pulse plethysmography relies on the fractional change in light absorption due to arterial pulsations [21] while MMSB relies on the magnetic disturbance due to blood flow.

From the waveform shown in Figure 40, it can be observed that the waveform acquired by the GMR sensor and the pulse plethysmography has similar periodicity with a difference in phase angle of approximately 90° . The phase difference is due to the inverted connection from the sensor output, V_{out+} and V_{out-} , to the input of the differential amplifier, $IN-$ and $IN+$, as shown in Figure 39. The inverted connection resulted in the inversion of the MMSB signal acquired from the GMR sensor.

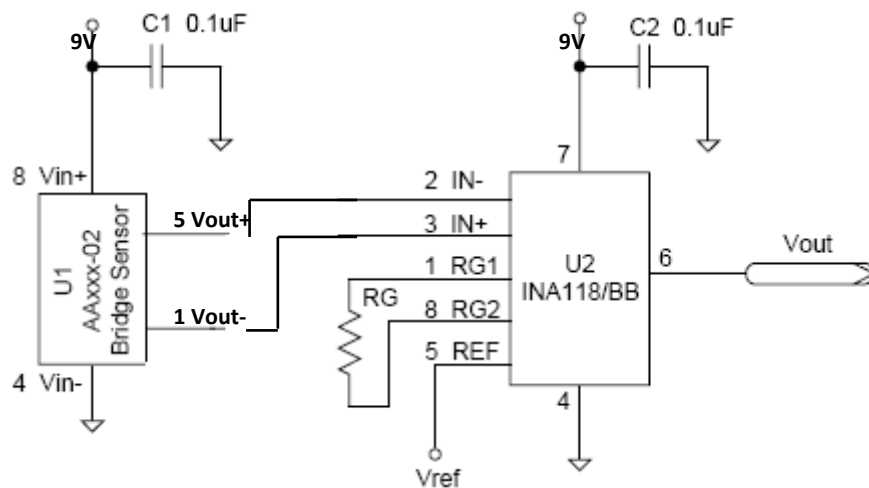


Figure 39 Amplifier circuit used for measurements of MMSB signal

Focusing on the phase difference between the MMSB and pulse plethysmography signal (refer to Figure 41), it was found that a time delay exist between the null of MMSB and the peak of the pulse plethysmography signal. In order to quantify the time delay, Figure 42 illustrates the results of the zoom-in plot of MMSB and pulse plethysmography signal.

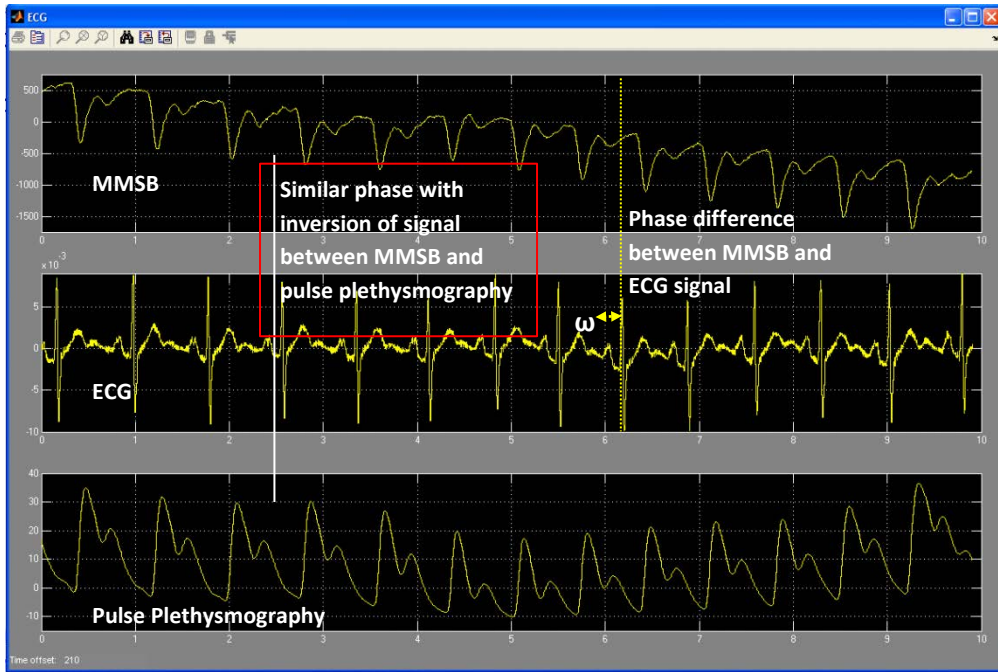


Figure 40 Illustration of MMSB signal collected together with ECG and pulse plethysmography using USBamplifier from Guger Technologies (gtec)

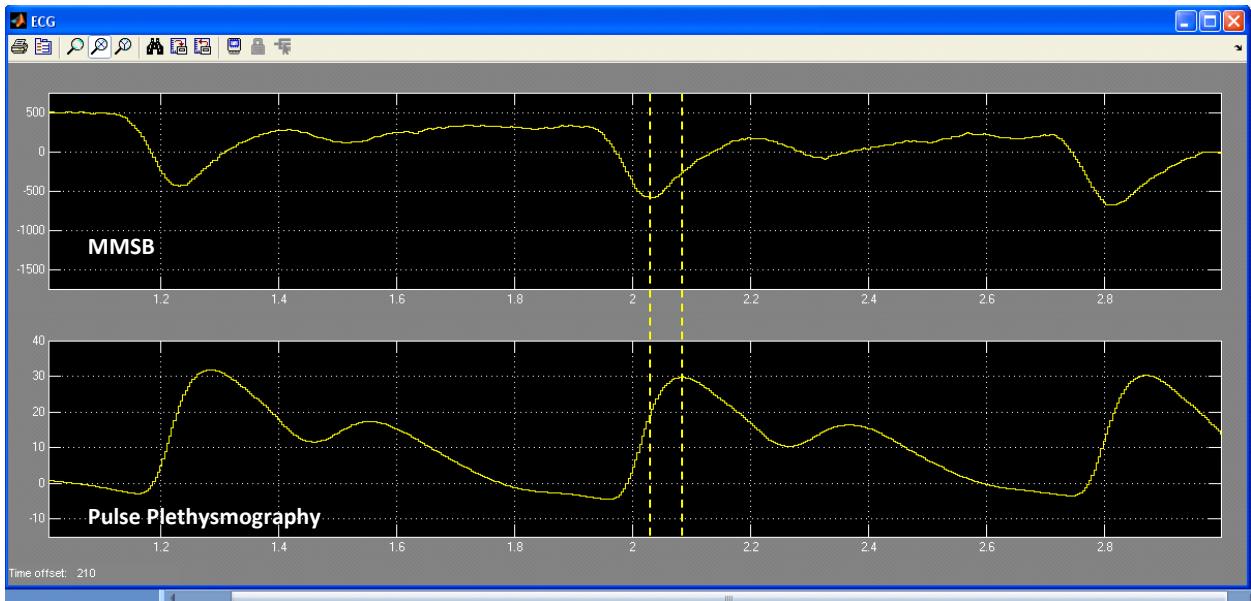


Figure 41 Illustration of MMSB signal collected together with pulse plethysmography using USBamplifier from Guger Technologies (gtec)

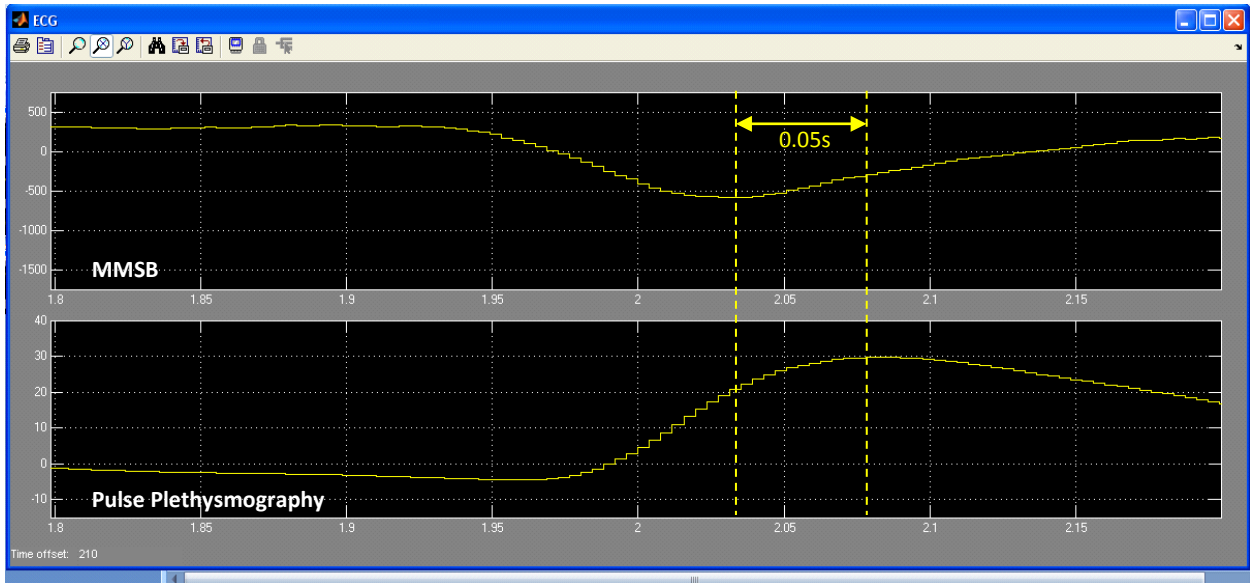


Figure 42 Zoom-in view of MMSB and pulse plethysmography signal acquired using the USBamplifier from Guger Technologies (gtec)

From Figure 42, it can be observed that a delay of 0.05 seconds exist between the two signals. This can be attributed to the difference in signal acquisition point where pulse plethysmography is acquired on the fingertip while MMSB is acquired on the wrist. The distance from the fingertip (pulse plethysmography) to the wrist (MMSB) was measured to be 24cm. Using the pulse wave velocity of 5m/s for healthy adult [28], the time delay from the fingertip (pulse plethysmography) to the wrist (MMSB) can be calculated as 0.048 seconds. As such, it can be concluded that the time delay between the MMSB and pulse plethysmography is due to the difference in location for signal acquisition.

In conclusion, this experimental setup has successfully demonstrated the accuracy of MMSB in acquiring the heart rate of the human subject with periodicity and phase comparison made to ECG and pulse plethysmography acquisition.

6 Comparison of measured waveform with Magnetocardiogram (MCG)

In this section, the waveform obtained in Figure 29 using the GMR sensor is compared with the MCG signal reported in [1]. Many studies of MCG have been made by mapping the vector content of the normal component of the magnetic field of the heart around the thorax. As the source of the MCG signal is the electric activity of the heart, the generation of the MCG signal from the progress of the activation front in the heart can be sketched with respect to the electrical activity of the heart as

shown in Figure 43. In addition, the x component of the MCG signal is also plotted as an illustration with respect to the heart activities as shown in Figure 43.

Using the magnetic vector generated by the MCG signal in the x, y and z component, the typical averaged magnetic heart vector during the QRS complex of the ECG signal can be plotted as shown in Figure 44. Using the vector average of the MCG_x , MCG_y and MCG_z signal, the vector MCG (VMCG) waveform can be extracted and plotted with the vector ECG (VECG) as shown in Figure 45 (a).

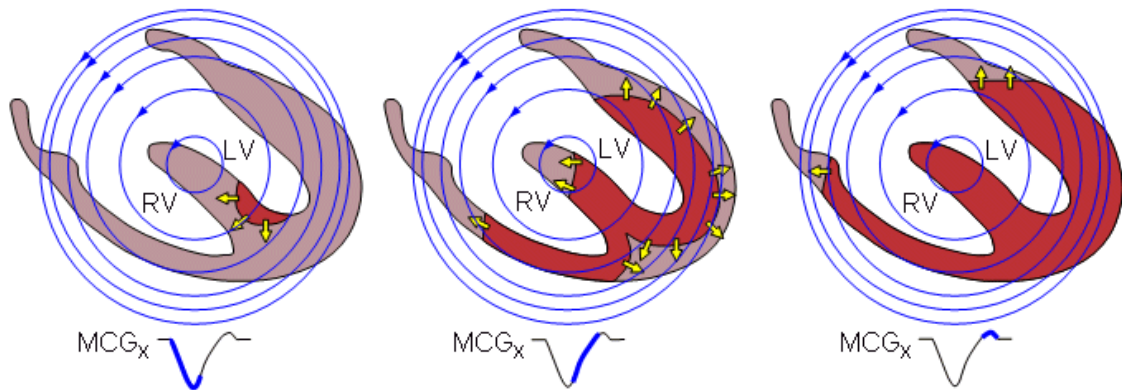


Figure 43 Illustration on the generation of the x component of the MCG signal with respect to the heart activities [1]

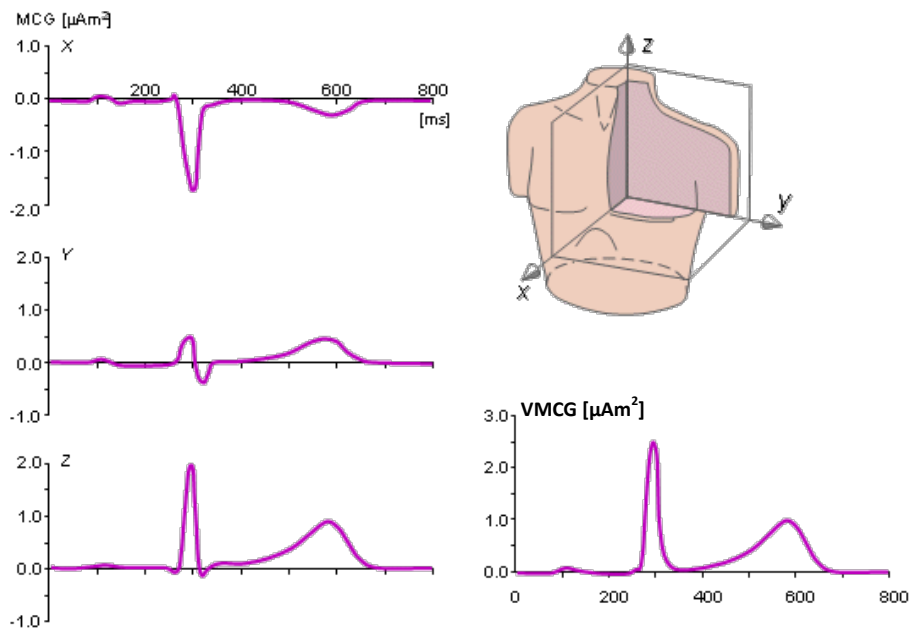


Figure 44 Vector MCG (VMCG) of a human subject obtained through the MCG_x , MCG_y and MCG_z vector plots acquired using the corrected uni-positional lead system

From Figure 45(a), it can be observed that the VMCG and VECG have an observable amount of phase difference during the QRS complex of the ECG signal and this had been reported [1] to vary from patient to patient and in various cases with cardiac disorders. The VMCG magnitude curve also

exhibits three different peaks during the QRS-complex of the electric heart vector: M_1 , M_2 and M_3 . M_1 appears to be generated by radial electric forces, M_2 by radial and tangential electric forces, and M_3 mainly by tangential electric forces.

Comparing the waveform characteristics acquired from the GMR sensor in Figure 45(b) with the peaks M_1 , M_2 and M_3 shown in Figure 45(a), similar peaks M_x , M_y and M_z can also be identified in Figure 45(b). Therefore, it can be concluded that the waveform acquired from the GMR sensor exhibits high correlations with the MHV magnitude curve during QRS-complex.

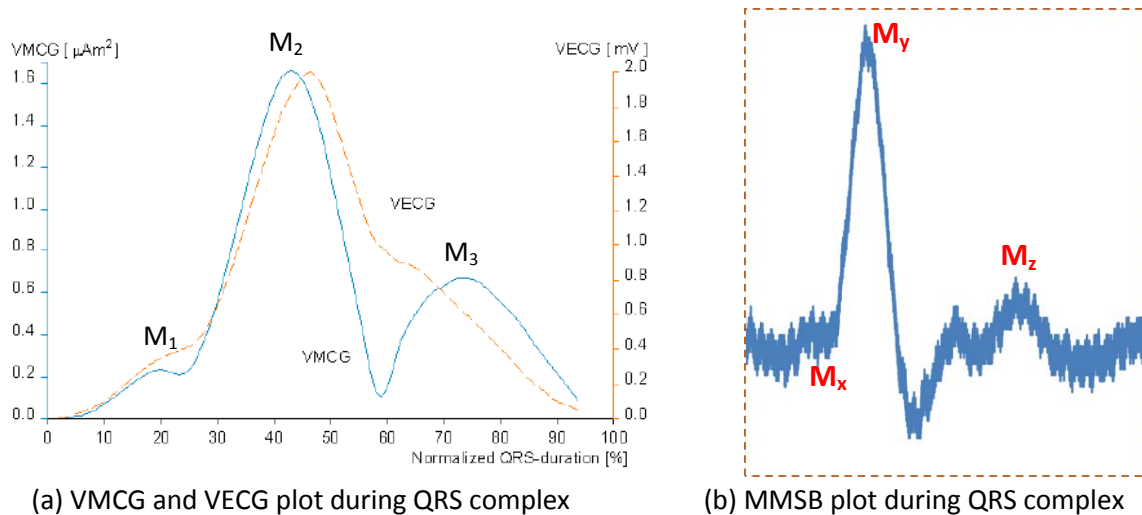


Figure 45 Simultaneous plots of the magnitude curves of the VECG (dashed curve) and the VMCG (solid curve) with MMSB plot during the QRS complex [1]

In addition to the comparison with the peaks in VMCG, on the basis of magnetic disturbance of the pulsatile blood flow on a uniform magnetic field, MMSB can be used to describe the activities of the heart as illustrated in Figure 46. For example, an increase in the measured magnetic disturbance can only be created by an increase in blood flow due to the compression of the heart ventricles. As the ventricles compress to their maximum, the peak of the waveform is reached. Without further mechanical force from the heart on the blood flow in the artery, the blood flow rate will reduce as illustrated in the decreasing amplitude of the waveform. With the relaxation of the ventricles, a back-flow of blood is present in the artery. Finally, the atrium will simultaneously compress, resulting in a small forward flow of blood as shown in the second peak detected in MMSB.

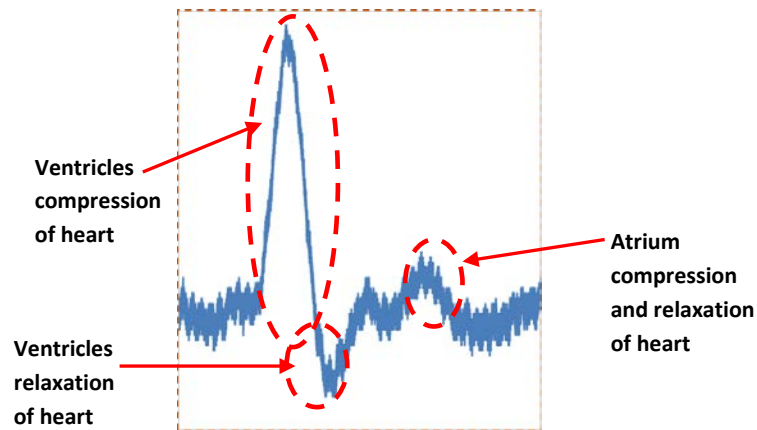


Figure 46 Illustration of MMSB and the activities related to the heart

7 Statistical validation of MMSB as a viable alternative to ECG

In this chapter, the heart rate (HR) obtained using MMSB will be statistically validated with respect to ECG, a gold standard instrumentation. This is because in clinical measurement, comparison of a new measurement technique with an established one is often needed to see whether they agree sufficiently for the new one (MMSB) to replace or supplement the existing one (ECG).

Four statistical tools typically used to validate such comparisons will be presented and discussed. These statistical tools are namely, Pearson Correlation and Coefficient of Determination [29], Paired Student's t-test [30]-[31], Wilcoxon Signed-Rank Test and Bland and Altman Test [35]-[36]. The results obtained from these statistical tests aims to support the use of MMSB as a viable alternative for measuring HR in resting condition for healthy individuals.

7.1 Measurement setup with data acquisition

In this section, the measurement setup with data acquisition for the concurrent HR acquisition using ECG and MMSB is shown in Figure 47. Waveform is acquired at a sampling rate of 1 KHz with resolution of 24 bits over +250mV range. Using this setup, measurements on 20 healthy subjects (aged 18-22) were conducted with 40 resting HR measurements collected.

Instantaneous HR is manually extracted from the time corresponding to two consecutive R peaks of ECG waveforms as illustrated in Figure 48(a). Using the MMSB waveforms, instantaneous HR is also manually extracted by locating the corresponding peaks (with respect to ECG R peaks) of two consecutive MMSB waveforms as shown in Figure 48(b). Instead of average HR, instantaneous HR is obtained from the waveforms of each subject so that the effect of averaging over multiple instantaneous HR will not affect the statistical validation of the data. The instantaneous HR extracted from ECG and MMSB pulses (as illustrated in Figure 48) are tabulated as shown in Table 4.

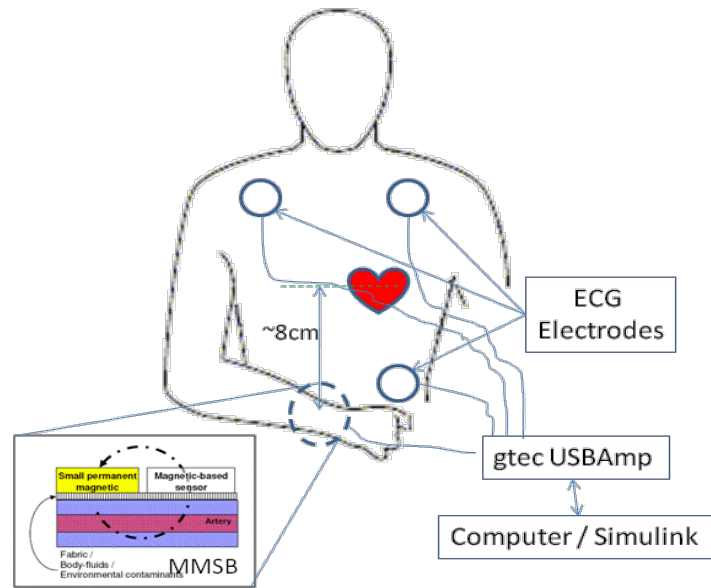
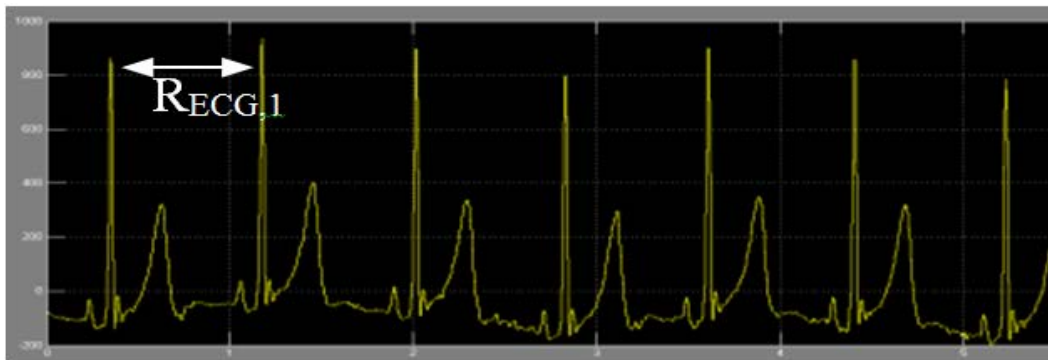
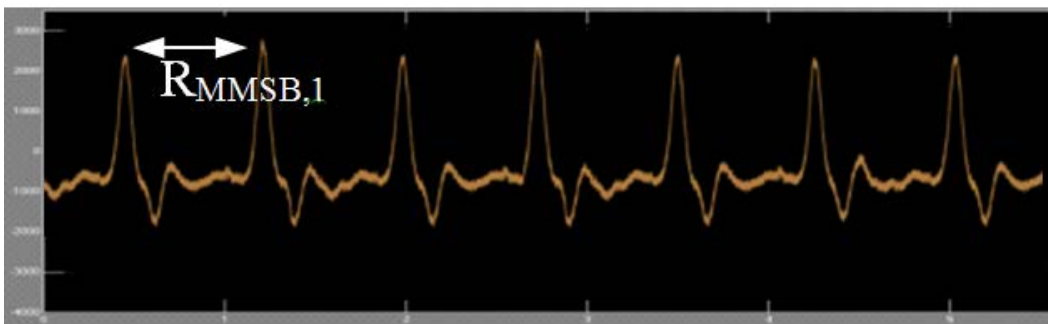


Figure 47 Concurrent HR acquisition using ECG and MMSB



(a) Extraction of instantaneous HR from ECG signal



(b) Extraction of instantaneous HR from MMSB signal

Figure 48 Extraction of instantaneous HR from ECG and MMSB waveforms

Measurement	Gold Standard (ECG)	MMSB Sensor	% w.r.t. ECG
1	78.174	78.271	-0.12
2	77.857	77.994	-0.18
3	78.174	78.694	-0.67
4	75.084	74.912	0.23
5	78.976	79.033	-0.07
6	78.094	78.311	-0.28
7	89.238	89.706	-0.52
8	95.557	95.654	-0.10
9	70.475	70.734	-0.37
10	76.540	76.747	-0.27
11	58.945	58.712	0.40
12	56.148	56.659	-0.91
13	83.160	82.810	0.42
14	93.071	92.973	0.11
15	77.426	77.251	0.23
16	73.121	73.253	-0.18
17	80.003	79.906	0.12
18	80.802	80.619	0.23
19	80.548	80.548	0.00
20	89.759	89.445	0.35

Measurement	Gold Standard (ECG)	MMSB Sensor	% w.r.t. ECG
21	80.170	80.674	-0.63
22	88.417	88.113	0.34
23	87.264	87.363	-0.11
24	92.918	92.838	0.09
25	84.998	84.783	0.25
26	100.553	100.160	0.39
27	75.047	74.966	0.11
28	80.911	80.911	0.00
29	80.488	80.488	0.00
30	71.390	71.390	0.00
31	91.888	92.215	-0.36
32	97.680	97.583	0.10
33	57.107	57.128	-0.04
34	57.982	58.055	-0.13
35	99.738	100.063	-0.33
36	93.809	94.038	-0.24
37	94.268	94.096	0.18
38	88.367	88.774	-0.46
39	65.746	65.793	-0.07
40	69.851	69.930	-0.11

Table 4 Instantaneous Heart Rate data collected for 40 measurements

7.2 Statistical validation - assessing agreement between two methods of measurement

7.2.1 Pearson Correlation and Coefficient of Determination

Pearson correlation coefficient (direction of relationship) and the Coefficient of Determination tests (strength of relationship) is used in the validation of the measurement outputs (i.e. instantaneous HR) of an alternative instrument (i.e. MMSB) against a “gold standard” (i.e. ECG) to ascertain the nature of relationship between two sets of scores [29].

The expression for Pearson’s Correlation Coefficient is shown in Equation 8.

$$\text{Equation 8} \quad r_{XY} = \left(\Sigma XY - ((\Sigma X)(\Sigma Y)/n) \right) / \left(\sqrt{(\Sigma X^2 - ((\Sigma X)^2/n))(\Sigma Y^2 - ((\Sigma Y)^2/n))} \right)$$

where r_{XY} = Pearson correlation coefficient

X = instantaneous HR for MMSB

Y = instantaneous HR for ECG

n = number of pairs of data (i.e. 40)

$$SS_X = \Sigma X^2 - ((\Sigma X)^2/n)$$

$$SS_Y = \Sigma Y^2 - ((\Sigma Y)^2/n)$$

Using the formula for Pearson’s Correlation Coefficient (reference Equation 8), r_{XY} and the Coefficient of Determination (r_{XY}^2) were calculated as shown in Table 5.

<i>Regression Statistics</i>	
Multiple R	0.99978
R Square	0.99956
Standard Deviation (SD)	0.24397
2xSD	0.48794

Table 5 Correlation test between Gold Reference and MMSB Sensor

From the results shown in Table 5, it can be observed that the HR data between MMSB and ECG are positively correlated (i.e. $r_{XY} = +0.9998$). The corresponding coefficient of determination (r_{XY}^2) is 0.9996. These results (i.e. $r_{XY}=0.9998$ and $r_{XY}^2 = 0.9998$) indicate that the relationship between the instantaneous HR measured by both the MMSB and ECG methods was almost perfectly linear^{***}.

^{***} If $r = 1.00$ there is a perfect positive relationship between the HR obtained from MMSB and ECG waveform (i.e. perfect match).

More importantly, the results highlight that < 1% of scores in the HR data from MMSB and ECG is due to error between the two measurements.

Using the instantaneous HR measured concurrently using the MMSB and an electrocardiogram system (g.USBamp, g.tec Medical Engineering GMBH, Austria), the instantaneous HR results are plotted as shown in Figure 49. To understand the direction of relationship, a linear regression is approximated based on the instantaneous HR results from the two measurements as shown in Figure 49. The linear equation obtained is shown in Equation 9 with regression ratio $R^2 = 0.9996$ obtained between Equation 9 and the data. The R^2 value being almost 1 indicates an almost perfect positive relationship between the measured data and the Equation 9.

Equation 9 $Y = 0.9971X + 0.2845$

where X = instantaneous HR for MMSB

Y = instantaneous HR for ECG

Based on the results obtained from this correlation study, it was concluded that the reproducibility of the HR measurement between the 2 devices is almost 1:1. The minute differences observable in this test was made possible through the resolution (i.e. up to 3 decimal places) provided by the sampling frequency of 1 KHz.

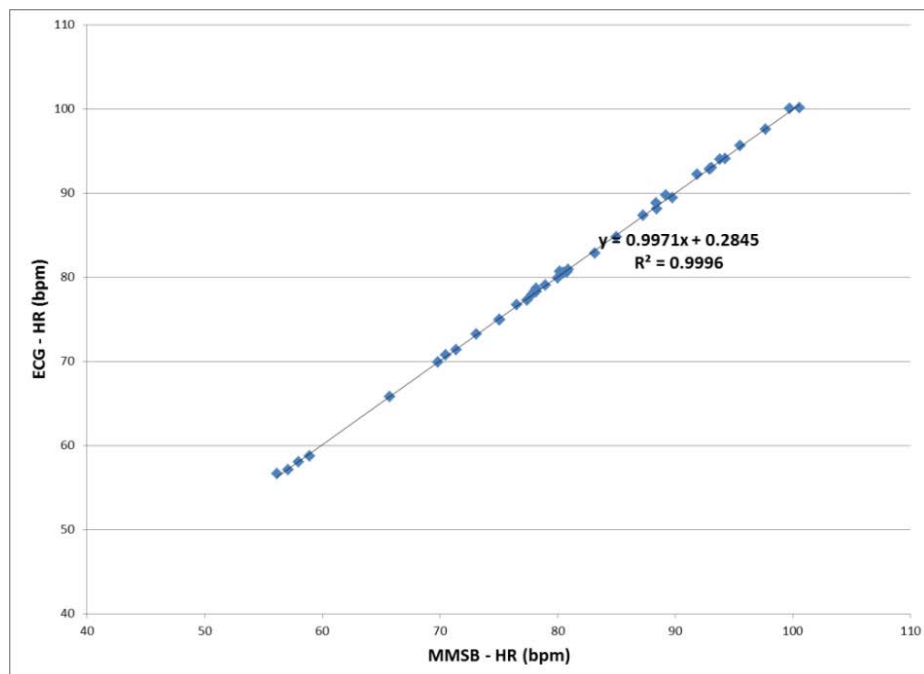


Figure 49 Plot of Correlation of HR obtained from MMSB and ECG waveforms in bpm⁺⁺⁺

⁺⁺⁺ Bpm = Beats per minute

7.2.2 Paired Student's t-test

The Paired Student's t-test is used as part of the scientific validation on the measurement outputs (refer to Table 4) to establish the statistical significance of the differences in the means between the two sets of scores [30].

From the instantaneous HR obtained in Table 4, the paired t-test determines whether HR measured from MMSB and ECG differ from each other in a significant way under the assumptions that the paired differences are independent and follows a normal distribution (i.e. $P^{***} < 0.05$).

The equation for obtaining the probability of obtaining the test statistic for the Paired Student t-test is shown in Equation 10.

$$\text{Equation 10} \quad P = (\bar{X} - \bar{Y}) / \sqrt{(s_1^2/n_1 + s_2^2/n_2)}$$

where P = probability of obtaining a test statistic

X = instantaneous HR obtained from MMSB

Y = instantaneous HR obtained from ECG

\bar{X} = mean of instantaneous HR obtained from MMSB

\bar{Y} = mean of instantaneous HR obtained from ECG

$n_1 = n_2$ = number of paired HR obtained from MMSB and ECG

s_1^2 = variance of instantaneous HR obtained from MMSB = $\Sigma(X_i - \bar{X})^2 / n_1$

s_2^2 = variance of instantaneous HR obtained from MMSB = $\Sigma(Y_i - \bar{Y})^2 / n_2$

The mean (i.e. \bar{X} and $\bar{Y} \approx 80$ bpm) and standard deviation (i.e. s_1 and s_2) were calculated and plotted as shown in Figure 50. Using Equation 10, the equation for Paired Student t-test, P was calculated as 0.24 (i.e. $P > 0.05$ which means the acceptance of the hypothesis). This result shows that there was no significant difference between the means of the HR measured by both devices (reference Figure 50).

*** P-value is the probability of obtaining a test statistic at least as extreme as the one that was actually observed, assuming that the null hypothesis is true. In this case a P-value less than 0.05 would result in the rejection of the null hypothesis at the 5% (significance) level. It means that if the HR is sampled on numerous occasions and interval estimates are made on each occasion, the resulting intervals would show the true HR parameter in approximately 95% of the cases. This is sufficient for typical clinical validation.

In addition, this result of the statistical test suggests the difference between the two sets of measurement is statistically not significant. Therefore, the results obtained from the Paired Student t-test supports the use of MMSB as a viable alternative to ECG for HR measurement.

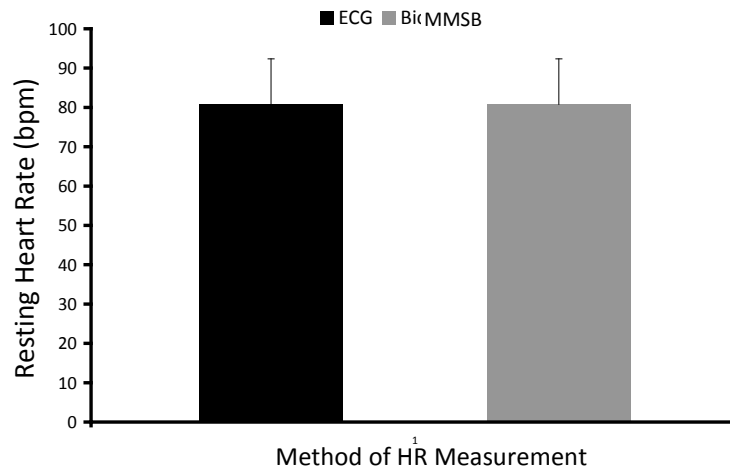


Figure 50 Mean and standard deviation of HR measured with the ECG and MMSB sensors (P = 0.24)

7.2.3 *Wilcoxon Matched-Pairs Signed-Ranks Test*

The Wilcoxon Signed-Rank Test is used as part of the scientific validation to determine whether there is a statistical significance on the difference in the median between paired samples [31][32].

The Wilcoxon Signed-Rank test was applied on the instantaneous HR measurement from Table 4

The Null Hypothesis for the Wilcoxon Signed-Rank Test (W) states that there was no difference between the paired samples at 95% limits of agreement (i.e. $P < 0.05$ for two-tailed non-directional test). The limit of agreement is selected to be similar to the Paired Student's t-test for consistency in the statistical validation of mean and median of HR measured.

In order to obtain the value of W, the difference between MMSB and ECG in Table 4 was calculated. These differences were ranked without regard to the sign of the difference (i.e., rank order the absolute differences). All zero differences were ignored (i.e., pairs with equal members). For differences with equal absolute values (i.e. ties), the same rank was assigned by taking the mean of the rank numbers that would had been assigned if they are different.

Upon completion of the ranking, the original signs on the differences were affix to the rank numbers. All positive ranks (W+) and all negative ranks (W-) were summed separately. At the same time, the total number of non-zero pairs (n) is recorded as shown in Table 6.

ECG	MMSB	Absolute value of difference	Rank	Rank if positive	Rank if negative
57.107	57.128	0.021	1		1
65.746	65.793	0.047	2		2
78.976	79.033	0.057	3		3
57.982	58.055	0.073	4		4
69.851	69.930	0.079	5		5
92.918	92.838	0.080	6	6	
75.047	74.966	0.081	7	7	
78.174	78.271	0.097	9		9
95.557	95.654	0.097	9		9
80.003	79.906	0.097	9	9	
97.680	97.583	0.097	11	11	
93.071	92.973	0.098	12	12	
87.264	87.363	0.099	13		13
73.121	73.253	0.132	14		14
77.857	77.994	0.137	15		15
75.084	74.912	0.172	16.5	16.5	
94.268	94.096	0.172	16.5	16.5	
77.426	77.251	0.175	18	18	
80.802	80.619	0.183	19	19	
76.54	76.747	0.207	20		20
84.998	84.783	0.215	21	21	
78.094	78.311	0.217	22		22
93.809	94.038	0.229	23		23
58.945	58.712	0.233	24	24	
70.475	70.734	0.259	25		25
88.417	88.113	0.304	26	26	
89.759	89.445	0.314	27	27	
99.738	100.063	0.325	28		28

ECG	MMSB	Absolute value of difference	Rank	Rank if positive	Rank if negative
91.888	92.215	0.327	29		29
83.160	82.810	0.350	30	30	
100.553	100.16	0.393	31	31	
88.367	88.774	0.407	32		32
89.238	89.706	0.468	33		33
80.170	80.674	0.504	34		34
56.148	56.659	0.511	35		35
78.174	78.694	0.520	36		36
80.548	80.548	0.000	-		
80.911	80.911	0.000	-		
80.488	80.488	0.000	-		
71.390	71.390	0.000	-		

Table 6 Illustration on quantification of ranking for Wilcoxon Matched-Pairs Signed-Ranks Test

From the sum of the positive and negative ranks obtained in Table 6, the variation of the median for the sum of the positive difference was $W+ = 274$ and the sum of the negative difference was $W- = 392$. The test statistic W was obtained from the minimum of $W+$ and $W-$ (i.e. $W = 274$).

The standard deviation Equation 11 for the measured pairs was used together with the mean, $\mu_w (=0$ for $n > 30$ [33]) for the Z_w score as shown in Equation 12.

Equation 11
$$\sigma_W = \sqrt{((n * (n + 1) * (2n + 1))/6)}$$

where σ_W = standard deviation for the sampling distribution of W (i.e. $\sigma_W = 127.30$)

n = number of pairs of non-zero difference between MMSB and ECG measurements (i.e. $n = 36$)

Equation 12
$$Z_W = ((W - \mu_W) \pm 0.5)/\sigma_W = ((W \pm 0.5)/\sigma_W)$$

where Z_W = standard deviation for the sampling distribution of W (i.e. $Z_w = 2.148, 2.141$)

To test for the Level of Significance, the table of critical values of Z_w (i.e. Table 7[33]) was used and it can be observed that the value of $Z_w = 2.148, 2.141$ was significant just a shade beyond the 0.025

level for a directional test. For a two-tailed non-directional test, it would be significant just beyond the 0.05 level (i.e. 95% limits of agreement).

Level of Significance for				
Directional Test				
0.05	0.025	0.01	0.005	0.0005
Non-Directional Test				
--	0.05	0.02	0.01	0.001
$Z_{w, \text{critical}}$				
1.645	1.960	2.326	2.576	3.291

Table 7 Critical Values of $\pm Z_w$ [33]

In conclusion, the Wilcoxon Signed-Rank test shows that there was no significant difference at 95% limits of agreement (i.e. two-tailed non-directional test) between the median of the HR measured by both MMSB and ECG methods. This result suggests that the difference between the two sets of measurement was statistically significant. As such, the Wilcoxon Signed-Rank test supports the use of MMSB as a viable alternative to ECG for HR measurement.

7.2.4 Bland and Altman Test

The Bland and Altman test (a.k.a. Limits of Agreement) is used in scientific validation on the measurement outputs of an alternative instrument (MMSB) against a “gold standard” (ECG) to determine the limits of agreement for the errors between the two sets of data. This analysis is recommended [35]-[36] when comparing the data between a “gold standard” and an alternative biomedical instrument.

The Bland and Altman method calculates the mean difference between MMSB and ECG methods of measurement (the ‘bias’), and 95% limits of agreement as the mean difference using Equation 13 (or 2*Standard Deviation - SD) [or more precisely (1.96*SD)]. It is expected that the 95% limits include 95% of differences between the two measurement methods.

Equation 13
$$SD = \sqrt{\Sigma(D_i - \bar{D})^2 / (N - 1)}$$

where D = difference between instantaneous HR obtained from MMSB and ECG

\bar{D} = mean of the difference between HR obtained from MMSB and ECG

N = total number of HR measurement samples (i.e. $n = 40$)

The plot obtained from this analysis is commonly called a Bland–Altman plot and the associated method is usually called the Bland–Altman method. Applying the Bland and Altman method on the measurements in Table 4, the Bland-Altman plot is shown in Figure 51. This analysis plots the difference (error) between the two HR measurements with respect to the means between the two scores.

Most of the errors should fall between ± 2 SD^{§§§} of the mean difference between the two HR measurement systems. Such a plot allows the assessment for systematic bias in the scores between the two systems. More importantly, the magnitude of the error at ± 2 SD is used to determine the clinical and practical significance of the measurement errors between the two systems. The magnitude of error should not be clinically or practically significant before the alternative device is accepted as a possible replacement for the Gold standard measurement device.

In the current analysis, the mean of the difference in HR measured between the ECG and the MMSB systems is -0.05 . The SD in HR measured between the ECG and the MMSB is ± 0.24 bpm (i.e. ± 0.48 bpm for ± 2 SD). The error data points are also evenly distributed across the range of heart rate measured, suggesting that there is no systematic error in the measurement. The ± 2 SD of 0.48 bpm is practically not significant for measurement of resting HR, suggesting that the MMSB system is a suitable alternative for the ECG system for HR measurement. For example, a resting HR of 70 bpm measured with the ECG system, would range between 69 bpm and 71 bpm in the HR measured by the MMSB system. Such a variation (± 1 bpm) is within the normal range of variation in HR measured in healthy individuals.

^{§§§} SD = Standard Deviation

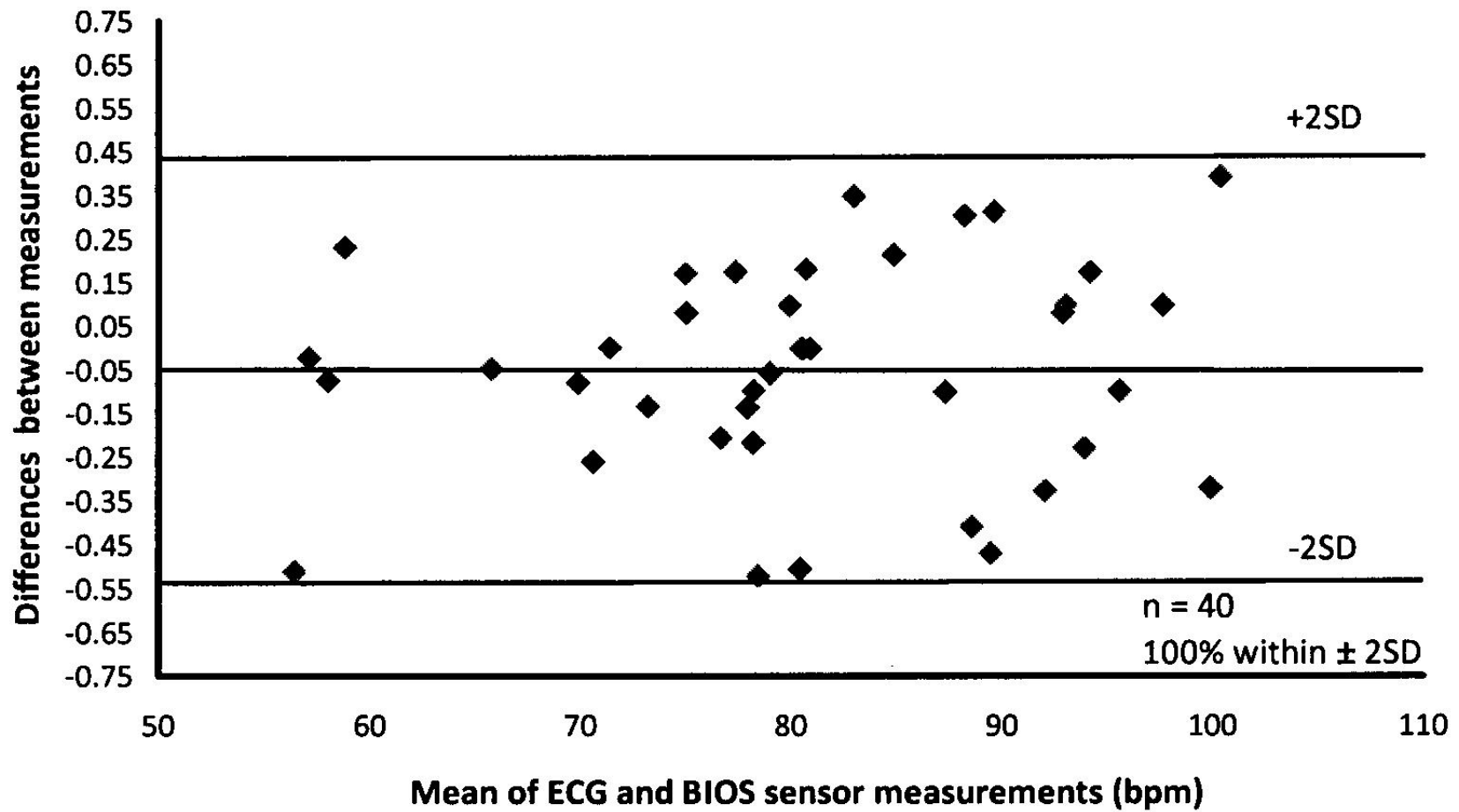


Figure 51 Plot of Bland and Altman test on HR obtained from MMSB and ECG waveforms

8 Conclusions

In this chapter, the experiment using Iron (II) Oxide (Fe_2O_3) fluidic flow has successfully demonstrated the presence of magnetic disturbance when a ferrofluid flows through a constant magnet field. This experiment provides the proof-of-concept for MMSB and forms the basis for further experiments on human subjects.

In addition, the experiments conducted on human subjects have successfully demonstrated the proof-of-operation of modulated magnetic signature of blood (MMSB) on human beings. Each of these experiment is designed to demonstrate the acquisition of heart rate using the magnetic disturbance measured through varying environmental conditions such as fabric, blood and underwater. Moreover, the waveform obtained using the GMR sensor is compared with ECG and pulse plethysmography where periodicity is concluded to be similar with known phase differences [27].

The waveform obtained using the GMR sensor was also compared to the MHV magnitude curve with good correlations on the waveform peaks related to the activities of the heart.

Finally the concurrent acquisition of HR using a gold standard (ECG) and MMSB on 20 subjects had been successfully completed with statistical analyses done. The results of the statistical analyses indicate that HR measured by the MMSB is almost perfectly correlated with HR measured with the ECG and there is a very good agreement between HR measured by the two devices. These data establish the use of the MMSB as a viable alternative to ECG in measuring HR healthy individuals under resting conditions.

Therefore, it is conclusive that MMSB is a feasible alternative to existing solutions of heart rate measurements with unique capability in sensing the pulse rate with direct electrical or optical contact with the human skin. Most important of all, the waveform acquired is proven to be accurate and reliable in measuring heart rate and qualifying its related activities.

Work in the next chapter of this dissertation will focus on the modelling of the MMSB phenomena in a multiphysics simulation software environment so as to allow the experimental setup to be optimised for implementation with high reliability and good sensitivity. In addition, the result obtained from the model will also support design size reduction allowing the sensing module (i.e. sensor and magnet) to be deployed as a wearable device.

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Chapter 3: Modelling and simulation of MMSB

1 Introduction

Current methods of heart or pulse rate acquisition have been classified in Chapter 1 into electrical [1][2], optical[3][7], microwave [1], acoustic [5][8][11], mechanical [6][9] or magnetic [10][12][13] means. Each of these methods has been studied in detail and its limitations have been well published and known.

In this chapter we will model first the experimental setup to demonstrate MMSB using iron oxide fluidic flow (described in Chapter 2, section 3.2). This is followed by the modelling of the magnetic method to acquire blood pulse using MMSB as proposed in this dissertation. In order to achieve these, the flow of the iron oxidic fluid and biomagnetic fluid (i.e. blood) in a weak magnetic field strength (0.1-0.2 Tesla) will be investigated.

With advancement in bioengineering and medical applications, extensive research work had been done and published on the fluid dynamics of biological fluids in the presence of magnetic field [14]-[16]. In order to examine the flow of a biomagnetic fluid under the action of an applied magnetic field, Haik et al [17] had developed a mathematical model for the Biomagnetic Fluid Dynamics (BFD), where the saturation or static magnetization is given by the Langevin magnetization equation. BFD differs from MagnetoHydroDynamics**** (MHD) in that BFD deals with no electric current and the fluidic flow is primarily affected by the magnetization of the fluid in the applied magnetic field.

In BFD, the behaviour of the biomagnetic fluid when it is exposed to magnetic field (magnetized) is described by the magnetization property M . Magnetization is the measure of how much the magnetic field is affecting the magnetic fluid and is a function of the magnetic field intensity H and the temperature T . However, in this chapter, the model will be developed with the assumption that the temperature of the system is a constant, where flow of blood in a weakly coupled magnetic field is affected purely by its magnetization properties.

**** MHD deals with the mathematical model of conducting fluids, which ignores the effect of polarization and magnetization.

2 Two-dimensional model of blood flow in the presence of a constant magnetic field

This model is based on the referred model [18] where bio-fluidic flow is coupled with magnetic particle flow (in this case, red blood cell) using the Arbitrary Lagrangian Eulerian (ALE) formulation. In this formulation, the governing equations for bio-fluidic flow are the continuity equation and the Navier-Stokes equations. For red blood cell movement, Newton's law is applied. The complete model includes the governing equations for the bio-fluidic flow, the governing equations for the motion of red blood cells, the interaction conditions between bio-fluidic and red blood cells at the interfaces, and boundary conditions.

To study the motion of red blood cell in bio-fluidic flow (i.e. blood flow), the red blood cells are assumed to be solid particles immersed in the bio-fluid or blood. In this case, the fluid-solid particle system occupies a bounded domain $\bar{\Omega}$ in \mathbb{R}^3 . At a typical instant of time t , Q particles occupy Q closed connected subsets $\sum_{q=1}^Q \Omega_q \subset \mathbb{R}^3$, which is surrounded by a viscous homogeneous fluid filling the domain $\bar{\Omega} - \sum_{q=1}^Q \Omega_q$ called the flow-channel area.

In this study the two coordinate systems used are:

1. A reference system, Ω , where the model is drawn and the particle movement is solved.
2. A moving mesh system, Ω_{def} , corresponding to the deformed mesh of the flow channel, where we simulate the fluid flow.

The time evolution of the domain Ω_{def} is determined by means of an Arbitrary Lagrangian-Eulerian (ALE) mapping $\mathbf{x}: \bar{\Omega} \times \mathbb{R}^3 \mapsto \Omega_{def}$ which maps any point (\mathbf{X}, t) to its image $\mathbf{x}(\mathbf{X}, t)$.

2.1 Transformation of coordinate systems

In the flow-channel area, the two coordinate systems, $(X, Y, Z) \in \Omega$ and $(x, y, z) \in \Omega_{def}$ are connected through a transformation T . At the initial state at $t = 0$, the two mesh systems are assumed to coincide. The transformation T maps the point initially located at (X, Y, Z) to the point (x, y, z) at time t as shown in Equation 14.

$$\text{Equation 14} \quad T: \begin{cases} x = x(X, Y, Z, t) \\ y = y(X, Y, Z, t) \\ z = z(X, Y, Z, t) \end{cases}$$

Solving Equation 14 at a particular instant of time, we can simplify Equation 14 to be independent of time t , as shown in Equation 15.

$$\text{Equation 15} \quad T: \begin{cases} x = x(X, Y, Z) \\ y = y(X, Y, Z) \\ z = z(X, Y, Z) \end{cases}$$

Suppose that the functions x , y , and z are continuous differentiable with respect to X , Y , Z . Then the infinitesimals dx , dy , dz transform of Equation 15 into dx , dy , dz according to Equation 16.

$$\text{Equation 16} \quad \begin{aligned} dx &= x_{,X}dX + x_{,Y}dY + x_{,Z}dZ \\ dy &= y_{,X}dX + y_{,Y}dY + y_{,Z}dZ \\ dz &= z_{,X}dX + z_{,Y}dY + z_{,Z}dZ \end{aligned}$$

where $x_{,X}$ denotes differentiation with respect to X .

Equation 16 can be written in the matrix form as shown in Equation 17 where the 3x3 matrix of partial derivatives is called the Jacobian matrix of the transformation, denoted by J in Equation 18.

$$\text{Equation 17} \quad \begin{bmatrix} dx \\ dy \\ dz \end{bmatrix} = \begin{bmatrix} x_{,X} & x_{,Y} & x_{,Z} \\ y_{,X} & y_{,Y} & y_{,Z} \\ z_{,X} & z_{,Y} & z_{,Z} \end{bmatrix} \begin{bmatrix} dX \\ dY \\ dZ \end{bmatrix}$$

$$\text{Equation 18} \quad |J| = x_{,X}(y_{,Y}z_{,Z} - y_{,Z}z_{,Y}) - x_{,Y}(y_{,X}z_{,Z} - y_{,Z}z_{,X}) + x_{,Z}(y_{,X}z_{,Y} - y_{,Y}z_{,X})$$

For $|J| \neq 0$, the transformation is invertible and there exists an inverse transformation at time t as shown in Equation 19.

$$\text{Equation 19} \quad T^{-1}: \begin{cases} X = X(x, y, z) \\ Y = Y(x, y, z) \\ Z = Z(x, y, z) \end{cases}$$

Re-writing Equation 19 in the matrix form, we have Equation 20, which can be denoted by J as shown in Equation 21.

$$\text{Equation 20} \quad \begin{bmatrix} dX \\ dY \\ dZ \end{bmatrix} = \begin{bmatrix} X_{,x} & X_{,y} & X_{,z} \\ Y_{,x} & Y_{,y} & Y_{,z} \\ Z_{,x} & Z_{,y} & Z_{,z} \end{bmatrix} \begin{bmatrix} dx \\ dy \\ dz \end{bmatrix}$$

$$\text{Equation 21} \quad \begin{bmatrix} dX \\ dY \\ dZ \end{bmatrix} = J^{-1} \begin{bmatrix} dx \\ dy \\ dz \end{bmatrix}$$

$$\text{where } J^{-1} = \begin{bmatrix} I_{Xx} & I_{Xy} & I_{Xz} \\ I_{Yx} & I_{Yy} & I_{Yz} \\ I_{Zx} & I_{Zy} & I_{Zz} \end{bmatrix} = \frac{1}{|J|} \mathbf{K},$$

$$K = \begin{bmatrix} y_{,Y}z_{,Z} - y_{,Z}z_{,Y} & x_{,Z}z_{,Y} - x_{,Y}z_{,Z} & x_{,Y}y_{,Z} - x_{,Z}y_{,Y} \\ y_{,Z}z_{,X} - y_{,X}z_{,Z} & x_{,X}z_{,Z} - x_{,Z}z_{,X} & y_{,X}x_{,Z} - y_{,Z}x_{,X} \\ y_{,X}z_{,Y} - y_{,Y}z_{,X} & x_{,Y}z_{,X} - x_{,X}z_{,Y} & x_{,X}y_{,Y} - x_{,Y}y_{,X} \end{bmatrix}$$

in which the \mathbf{I} syntax is used to highlight the computation of the Jacobian of the inverse transformation from the inverse of the original Jacobian.

Equating Equation 20 and Equation 21, we have the relations described in Equation 22, which are crucial in transforming the calculation results from Ω_{def} to Ω .

$$\begin{aligned} X_{.x} &= (y_{,Y}z_{,Z} - y_{,Z}z_{,Y})/|J| \\ X_{.y} &= (x_{,Z}z_{,Y} - x_{,Y}z_{,Z})/|J| \\ X_{.z} &= (x_{,Y}y_{,Z} - x_{,Z}y_{,Y})/|J| \\ \\ Y_{.x} &= (y_{,Z}z_{,X} - y_{,X}z_{,Z})/|J| \\ Y_{.y} &= (x_{,X}z_{,Z} - x_{,Z}z_{,X})/|J| \\ Y_{.z} &= (y_{,X}x_{,Z} - y_{,Z}x_{,X})/|J| \\ \\ Z_{.x} &= (y_{,X}z_{,Y} - y_{,Y}z_{,X})/|J| \\ Z_{.y} &= (x_{,Y}z_{,X} - x_{,X}z_{,Y})/|J| \\ Z_{.z} &= (x_{,X}y_{,Y} - x_{,Y}y_{,X})/|J| \end{aligned}$$

Equation 22

These relations are used in the software COMSOL for the calculation of numerical results through the transforming from Ω_{def} to Ω .

2.2 Motion of Fluid-Solid Flow in the Deformed Mesh System

To study the motion of red blood cell in the blood vessels (i.e fluid flow channel), we assume that the gravitational force can be neglected and the red blood cell movement is governed by Newton's second law as shown in Equation 23.

$$\begin{aligned} m_q \frac{\partial V_q}{\partial t} &= F_v + F_q + F_{\text{mag}}, & q &= 1, 2, 3, \dots, Q \\ V_q|_{t=0} &= 0, & & \text{initial conditions} \end{aligned}$$

Equation 23

where V_q = velocity vector of the qth red blood cell
 m_q = mass of the qth red blood cell
 F_v = drag force
 F_q = collision force
 F_{mag} = magnetic force

The three applied loads, drag force F_v , collision force F_q , and magnetic force F_{mag} , are defined based on the following assumptions:

1. All boundaries of particles experience drag force F_v from fluid as shown in Equation 24.

$$\text{Equation 24} \quad \mathbf{F}_v = -n_f \cdot \left(-p\mathbf{I} + \eta(\nabla\mathbf{v} + (\nabla\mathbf{v})^T) \right)$$

where n_f = viscous drag of the fluid

η = dynamic viscosity

p = the pressure

\mathbf{v} = velocity vector

\mathbf{I} = 3×3 identity matrix

2. To prevent the collisions among the red blood cell, and between the red blood cells and the vessel walls, the red blood cell to red blood cell interaction force $F_{q,p}$ and the red blood cell-wall interaction force $F_{q,w}$ are applied when the distance between two red blood cells, or between a red blood cell and a wall, is within the order of the element size [19]. These interaction forces are illustrated in Figure 52 and described mathematically in Equation 25 to Equation 27.

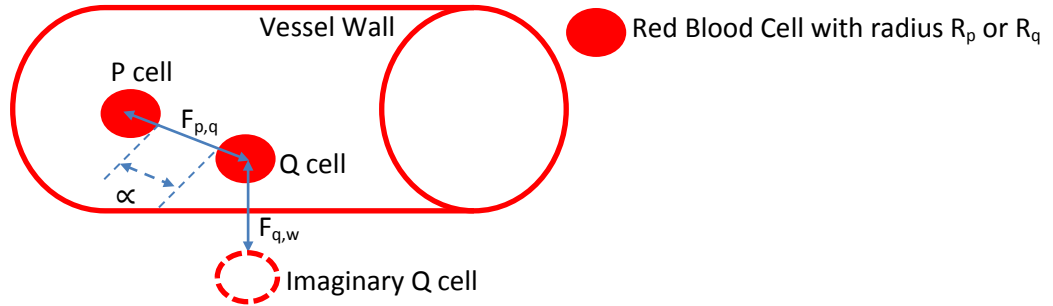


Figure 52 Illustration of the interaction forces between red blood cells and the vessel walls

$$\text{Equation 25} \quad \mathbf{F}_q = \sum_{p=1, p \neq q}^0 \mathbf{F}_{q,p} + \sum_{w=1}^2 \mathbf{F}_{q,w}$$

in which,

$$\text{Equation 26} \quad \mathbf{F}_{q,p} = \begin{cases} \mathbf{0}, & \text{for } d_{q,p} > R_q + R_p + \alpha \\ \frac{1}{\varepsilon_q} (\mathbf{X}_q - \mathbf{X}_p) (R_q + R_p + \alpha - d_{q,p}), & \text{for } d_{q,p} \leq R_q + R_p + \alpha \end{cases}$$

and

$$\text{Equation 27} \quad \mathbf{F}_{q,w} = \begin{cases} \mathbf{0}, & \text{for } d_{q,w} > 2R_q + \alpha \\ \frac{1}{\varepsilon_w} (\mathbf{X}_q - \mathbf{X}_w) (2R_q + \alpha - d_{q,p}), & \text{for } d_{q,p} \leq 2R_q + \alpha \end{cases}$$

where $d_{q,p}$ denotes the distance between the centres of the q th and p th red blood cell, $d_{q,w}$ denotes the distance between the centres of the q th red blood cell and the imaginary red blood cell on the other side of the wall, X_q and R_q are centre and radius of the q th red blood cell, α is the distance between the p th and q th red blood cell where the force is non-zero, and ϵ_q and ϵ_w are small positive stiffness parameters of the particle and wall respectively.

3. To model the magnetic disturbance of the blood at the measurement site, an external magnetic field is applied to generate magnetic force acting on the red blood cell. This force, which is composed of 3 components, is governed by the equations:

Equation 28
$$\mathbf{F}_{mag} = \frac{1}{\mu_r} (\mathbf{M} \cdot \nabla) \mathbf{B}$$

where μ_r is the relative permeability of the red blood cell

$\mathbf{M} = (M_x, M_y, M_z)$ is the magnetic moment of the red blood cell

$\mathbf{B} = (B_x, B_y, B_z)$ is the magnetic flux density in 3-dimensional space

To determine the drag force F_v in Equation 24, blood is assumed to be an isotropic, homogeneous incompressible fluid. The motion of the blood is described by the continuity Equation 29 and the Navier-Stokes Equation 30.

Equation 29
$$\nabla \cdot \mathbf{v} = 0$$

Equation 30
$$\rho_f \frac{\partial \mathbf{v}}{\partial t} + \rho_f (\mathbf{v} \cdot \nabla) \mathbf{v} - \nabla \cdot \boldsymbol{\sigma} = \mathbf{F}$$

where ρ_f denotes the blood density

$\mathbf{v} = [u, v, w]^T$ represents the 3D velocity vector

\mathbf{F} is the volume force acting on the fluid

For this model, we neglect the effect of gravitational force and thus, $\mathbf{F} = 0$. The quantity $\boldsymbol{\sigma}$ in Equation 30 is the stress tensor given by Equation 31.

Equation 31
$$\boldsymbol{\sigma} = -p\mathbf{I} + \eta(\nabla \mathbf{v} + (\nabla \mathbf{v})^T)$$

where η = blood viscosity

\mathbf{v} = velocity vector

p = blood pressure

\mathbf{I} = 3x3 identity matrix

On the wall, the no-slip condition is applied. On the inflow boundary Γ_{in} , the velocity is assumed to be constant, whereas on the outflow boundary Γ_{out} , the stress-free condition as described by Equation 32.

$$\begin{aligned} \text{Equation 32} \quad & \mathbf{v} = \mathbf{v}_0 \text{ on } \Gamma_{in} \\ & \boldsymbol{\sigma} \cdot \mathbf{n} = \mathbf{0} \text{ on } \Gamma_{out} \end{aligned}$$

For static condition in stationary bodies, the magnetic flux density \mathbf{B} is governed by the Maxwell-Ampere's law for the magnetic field \mathbf{H} (A/m) and the current density \mathbf{J} (A/m²) in Equation 33 can be applied (where $\mathbf{J} = 0$ for non-conducting fluid). In addition, using the Gauss' law for the magnetic flux density \mathbf{B} (Vs/m²) stated in Equation 34, this will result in the constitutive equations describing the relationships between \mathbf{B} and \mathbf{H} in the different parts of the modelling domain in Equation 35.

$$\text{Equation 33} \quad \nabla \times \mathbf{H} = \mathbf{J} = \mathbf{0}$$

$$\text{Equation 34} \quad \nabla \times \mathbf{B} = \mathbf{0}$$

$$\text{Equation 35} \quad \mathbf{B} = \begin{cases} \mu_0 \mu_{r,mag} \mathbf{H} + \mathbf{B}_{rem} & \text{permanent magnet} \\ \mu_0 (\mathbf{H} + \mathbf{M}_{ff}(\mathbf{H})) & \text{red blood cell in blood stream} \\ \mu_0 \mathbf{H} & \text{tissue and air} \end{cases}$$

where μ_0 = magnetic permeability of vacuum (Vs/(A·m))

$\mu_{r,mag}$ = relative magnetic permeability of the permanent magnet

\mathbf{B}_{rem} = residual magnetic flux (A/m)

\mathbf{M}_{ff} = magnetization vector^{****} in the blood stream (A/m), which is a function of magnetic field, \mathbf{H}

Defining a magnetic vector potential \mathbf{A} such that $\mathbf{B} = \nabla \times \mathbf{A}$, $\nabla \cdot \mathbf{A} = 0$ and substituting this into Equation 33 through Equation 35, the equations can be simplified to a 2D problem with no perpendicular currents as shown in Equation 36.

$$\text{Equation 36} \quad \nabla \times \left(\frac{1}{\mu_0} \nabla \times \mathbf{A} - \mathbf{M}_{ff} \right) = \mathbf{J}$$

where $\mathbf{J} = 0$ for 2D with no perpendicular currents

Using the identity in Equation 37 and the Coulomb gauge $\nabla \cdot \mathbf{A} = 0$, the Equation 36 takes the form as shown in Equation 38.

$$\text{Equation 37} \quad \nabla \times (\nabla \times \mathbf{A}) = \nabla(\nabla \cdot \mathbf{A}) - \nabla^2 \mathbf{A}$$

^{****} Magnetization is the measure of how much the magnetic field affects the magnetic fluid. In this case, it is the behaviour of the biomagnetic fluid (blood) when it is exposed to magnetic field (magnetized).

Equation 38
$$\nabla A = -\nabla \times (\mu_0 \mathbf{M}_{ff})$$

Due to the movement of the coordinate system, the mesh velocity $\Psi = (\Psi_x; \Psi_y; \Psi_z)$ is introduced in the deformed domain Ω_{def} , as explained in the introduction on page 64. To guarantee a smoothly varying distribution of the nodes, we assume that the nodes on $\partial\Omega_q$ move with the red blood cell (no slip) and that each component of the mesh velocity in the fluid channel is governed by a Laplace equation as shown in Equation 39.

Equation 39
$$\nabla^2 \Psi = \mathbf{0}, \forall \mathbf{x} \in \Omega_{def}$$

The above equation is used to smooth gradient of the mesh velocity over the domain so as to reduce mesh distortion. Once the mesh velocity components are determined, we can determine the smoothed deformed mesh for the flow channel at each time instant by updating the coordinates of the nodes according to the following formulae:

Equation 40
$$\mathbf{x} = \mathbf{X} + \int_0^t \Psi_x dt$$

Equation 41
$$\mathbf{y} = \mathbf{Y} + \int_0^t \Psi_y dt$$

Equation 42
$$\mathbf{z} = \mathbf{Z} + \int_0^t \Psi_z dt$$

Another condition that needs to be specified is that the fluid, red blood cell and mesh move with the same velocity on the particle boundaries, i.e.,

Equation 43
$$\Psi = \mathbf{v} = \mathbf{V}_q \text{ on } \partial\Omega_q$$

We now have the strong coupled problem for the fluid-red blood cell flow in the blood circulatory system. These equations are solved to yield \mathbf{V}_q in Ω and the velocity \mathbf{v} , the pressure p , the magnetic potential A and the mesh velocity Ψ , in Ω_{def} .

The formulation of the fluid flow problem is to find $(\mathbf{v}, p, A, \Psi) \in \mathfrak{S}$ in the deformed mesh system at each time instant such that all the Dirichlet Boundary conditions are satisfied and $\forall (\hat{\mathbf{v}}, \hat{\mathbf{p}}, \hat{\mathbf{A}}, \hat{\Psi}) \in \mathfrak{S}^0 \equiv \{(\hat{\mathbf{v}}, \hat{\mathbf{p}}, \hat{\mathbf{A}}, \hat{\Psi}) \in \mathfrak{S}^0 | \hat{\mathbf{v}} = \mathbf{0} \text{ on } \partial\Omega_{def}, \hat{\mathbf{p}} = \mathbf{0} \text{ on } \partial\Omega_{def}, \hat{\mathbf{A}} = \mathbf{0} \text{ on } \partial\Omega_{def}, \hat{\Psi} = \mathbf{0} \text{ on } \partial\Omega_{def}\}$.

Equation 44
$$\int_{\Omega_{def_v}} \hat{\mathbf{p}}(\nabla \cdot \mathbf{v}) d\Omega = \mathbf{0}$$

Equation 45
$$\int_{\Omega_{def_\rho}} \left(\rho_f \hat{\mathbf{v}} \cdot \frac{\partial \mathbf{v}}{\partial t} + \eta \nabla \hat{\mathbf{v}} : \nabla \mathbf{v} + \rho_f \hat{\mathbf{v}} \cdot (\mathbf{v} \cdot \nabla) \mathbf{v} - \rho \nabla \cdot \hat{\mathbf{v}} \right) d\Omega = \int_{\partial\Omega_{def}} \hat{\mathbf{v}}(\boldsymbol{\sigma} \cdot \mathbf{n}) ds$$

Equation 46
$$\int_{\Omega_{def_A}} \left(\nabla \hat{\mathbf{A}} : \nabla \mathbf{A} - \hat{\mathbf{A}} \cdot \nabla \times (\mu_0 \mathbf{M}_{ff}) \right) d\Omega = \mathbf{0}$$

Equation 47
$$\int_{\Omega_{def_\Psi}} (\nabla \hat{\Psi} : \nabla \Psi) d\Omega = \mathbf{0}$$

where Ω_{def_v} , $\partial\Omega_{def_p}$, $\partial\Omega_{def_A}$ and $\partial\Omega_{def_\Psi}$ are the parts of boundary where the velocity, the pressure, the magnetic potential and the mesh velocity are specified. It should also be addressed that various surface integral terms, arising in the formulation, vanish as the test functions involved in the terms in are zero on the boundary.

Since the computations are conducted in the reference coordinates, Ω , we need to transform Equation 43 - Equation 47 in the deformed coordinates to those equations in the reference coordinates. Through this and using Equation 32, we will obtain Equation 48- Equation 51.

$$\text{Equation 48} \quad \int_{\Omega} \hat{p}(\nabla \cdot \mathbf{v}) |J| d\Omega = 0$$

$$\text{Equation 49} \quad \int_{\Omega} \left(\rho_f \hat{\mathbf{v}} \cdot \frac{\partial \mathbf{v}}{\partial t} + \eta \nabla \hat{\mathbf{v}} : \nabla \mathbf{v} + \rho_f \hat{\mathbf{v}} \cdot (\mathbf{v} \cdot \nabla) \mathbf{v} - \rho \nabla \cdot \hat{\mathbf{v}} \right) |J| d\Omega = 0$$

$$\text{Equation 50} \quad \int_{\Omega} \left(\nabla \hat{\mathbf{A}} : \nabla \mathbf{A} - \hat{\mathbf{A}} \cdot \nabla \times (\mu_0 \mathbf{M}_{ff}) \right) |J| d\Omega = 0$$

$$\text{Equation 51} \quad \int_{\Omega_{def}} (\nabla \hat{\Psi} : \nabla \Psi) |J| d\Omega = 0$$

where the derivatives of the unknown functions $\Psi_i (i = x, y, z)$ are determined by the following expressions:

$$\begin{aligned} \Psi_{i,x} &= \Psi_{i,X} I_{X,x} + \Psi_{i,Y} I_{Y,x} + \Psi_{i,Z} I_{Z,x} \\ \Psi_{i,y} &= \Psi_{i,X} I_{X,y} + \Psi_{i,Y} I_{Y,y} + \Psi_{i,Z} I_{Z,y} \\ \Psi_{i,z} &= \Psi_{i,X} I_{X,z} + \Psi_{i,Y} I_{Y,z} + \Psi_{i,Z} I_{Z,z} \end{aligned}$$

Equation 52

and for the test functions:

$$\begin{aligned} \hat{\Psi}_{i,x} &= \hat{\Psi}_{i,X} I_{X,x} + \hat{\Psi}_{i,Y} I_{Y,x} + \hat{\Psi}_{i,Z} I_{Z,x} \\ \hat{\Psi}_{i,y} &= \hat{\Psi}_{i,X} I_{X,y} + \hat{\Psi}_{i,Y} I_{Y,y} + \hat{\Psi}_{i,Z} I_{Z,y} \\ \hat{\Psi}_{i,z} &= \hat{\Psi}_{i,X} I_{X,z} + \hat{\Psi}_{i,Y} I_{Y,z} + \hat{\Psi}_{i,Z} I_{Z,z} \end{aligned}$$

Equation 53

The derivatives of other unknown functions u , v , w , A_x , A_y and A_z are defined in the same way as those of the Ψ_i functions.

2.3 Numerical results using multiphysics simulation software

In a two dimension case, the magnetic potential is assumed to have a nonzero component only in the direction A_z , perpendicular to the (x,y) plane (i.e. $\mathbf{A} = (0, 0, A_z)$). On $\partial\Omega_q$ and $\partial\Omega$, the magnetic potential is set to zero, that is, $A_z = 0$.

As such, the magnetization vector in the blood stream can be expressed as shown in Equation 54.

Equation 54
$$M_{ffx} = \frac{1}{\mu_0} \frac{\partial A_z}{\partial y}, \quad M_{ffy} = \frac{1}{\mu_0} \frac{\partial A_z}{\partial x}$$

Based on the results presented [41], the magnetization curves for a soft magnetic material^{****}, can be approximated by simple two-parameter arctangent functions of the form $M = \alpha \cdot \arctan(\beta \cdot M_{ff})$, where the arctangent expression includes two material parameters α (A/m), saturation magnetization, and β (m/A). Using the notation of magnetization in the (x,y) axis as M_x and M_y respectively, the induced magnetization of blood can be expressed as shown in Equation 55.

Equation 55
$$M_x = \alpha \tan^{-1} \left(\frac{\beta}{\mu_0} \frac{\partial A_z}{\partial y} \right), \quad M_y = \alpha \tan^{-1} \left(\frac{\beta}{\mu_0} \frac{\partial A_z}{\partial x} \right)$$

Linearizing Equation 55 for analysis of magnetization in weak magnetic field where $\frac{\beta}{\mu_0} \frac{\partial A_z}{\partial x}$ and $\frac{\beta}{\mu_0} \frac{\partial A_z}{\partial y}$ are numerically small (i.e. $\ll 1$), the arctangent function in Equation 55 can be simplified as Equation 56.

Equation 56
$$M_x = \frac{\chi}{\mu_0} \frac{\partial A_z}{\partial y}, \quad M_y = \frac{\chi}{\mu_0} \frac{\partial A_z}{\partial x}$$

where $\chi = \alpha\beta$ is the magnetic susceptibility

With the assumption that the red blood cells in the blood fluid do not interact, the magnetic force $F = (F_x, F_y)$ on the blood for relatively weak fields is given by $\vec{F} = (\vec{m} \cdot \vec{\nabla}) \vec{H}$ [42] which can be simplified to a 2D problem with no perpendicular currents. As such, for a 2D problem, the operator $\vec{\nabla}$ can be simplified as $\vec{\nabla} = \tilde{i} \frac{\partial}{\partial x} + \tilde{j} \frac{\partial}{\partial y}$ and the magnetic flux density \vec{B} can be expressed as $\vec{H} = \frac{1}{\mu_0 \mu_r} \vec{B} = \frac{1}{\mu_0 \mu_r} \left[\tilde{i} \frac{\partial A_z}{\partial y} - \tilde{j} \frac{\partial A_z}{\partial x} \right]$.

Re-writing the results in Equation 56 as a vector form, $\vec{m} = \tilde{i} \frac{\chi}{\mu_0} \frac{\partial A_z}{\partial y} + \tilde{j} \frac{\chi}{\mu_0} \frac{\partial A_z}{\partial x}$.

Therefore,
$$\vec{F} = (\vec{m} \cdot \vec{\nabla}) \vec{H}$$

$$\begin{aligned} \vec{F} &= \left(\left[\tilde{i} \frac{\chi}{\mu_0} \frac{\partial A_z}{\partial y} + \tilde{j} \frac{\chi}{\mu_0} \frac{\partial A_z}{\partial x} \right] \cdot \left[\tilde{i} \frac{\partial}{\partial x} + \tilde{j} \frac{\partial}{\partial y} \right] \right) \left[\tilde{i} \frac{\partial A_z}{\partial y} - \tilde{j} \frac{\partial A_z}{\partial x} \right] \frac{1}{\mu_0 \mu_r} \\ &= \tilde{i} \left(\frac{\chi}{\mu_r \mu_0^2} \frac{\partial A_z}{\partial y} \frac{\partial^2 A_z}{\partial y \partial x} + \frac{\chi}{\mu_r \mu_0^2} \frac{\partial A_z}{\partial x} \frac{\partial^2 A_z}{\partial y^2} \right) - \tilde{j} \left(\frac{\chi}{\mu_r \mu_0^2} \frac{\partial A_z}{\partial y} \frac{\partial^2 A_z}{\partial x^2} + \frac{\chi}{\mu_r \mu_0^2} \frac{\partial A_z}{\partial x} \frac{\partial^2 A_z}{\partial x \partial y} \right) \end{aligned}$$

Equation 57
$$F_x = \frac{\chi}{\mu_r \mu_0^2} \left(\frac{\partial A_z}{\partial x} \frac{\partial^2 A_z}{\partial y^2} + \frac{\partial A_z}{\partial y} \frac{\partial^2 A_z}{\partial y \partial x} \right), \quad F_y = \frac{\chi}{\mu_r \mu_0^2} \left(\frac{\partial A_z}{\partial x} \frac{\partial^2 A_z}{\partial x \partial y} + \frac{\partial A_z}{\partial y} \frac{\partial^2 A_z}{\partial x^2} \right)$$

^{****} Material that attains magnetization easily and reaches saturation at relatively low magnetic field. At the same time, loses magnetization quickly as the external field diminishes.

The final expression for the volume force in the blood stream can be obtained by multiplying Equation 57^{§§§§} by the ferrofluid mass fraction, K_{ff} . This will result in the expressions shown in Table 8 allowing the magnetic field at the sensor location to be solved in COMSOL environment.

The solved magnetic field will be translated to voltage output based on the sensor response characteristics as shown in Table 11, which will be verified with experimental measurements at latter part of this chapter.

Name	Expression	Description
Mu0	Mu0_qa	Permeability of vacuum
M_ffx	$k_{ff} * (\chi_{ff} / \mu_0) * A_{zy}$	Induced ferrofluid magnetization, x-component
M_ffy	$-k_{ff} * (\chi_{ff} / \mu_0) * A_{zx}$	Induced ferrofluid magnetization, y-component
F_ffx	$k_{ff} * (A_{zx} * A_{zyy} + A_{zy} * A_{zxy}) * \chi_{ff} / ((\mu_0)^2 * (1 + \chi_{ff}))$	Ferrofluid volume force, x-component
F_ffy	$k_{ff} * (A_{zx} * A_{zxy} + A_{zy} * A_{zxx}) * \chi_{ff} / ((\mu_0)^2 * (1 + \chi_{ff}))$	Ferrofluid volume force, y-component

where notations in COMSOL are $A_{zy} = \frac{\partial A_z}{\partial y}$, $A_{zx} = \frac{\partial A_z}{\partial x}$, $A_{zxx} = \frac{\partial A_z}{\partial x^2}$, $A_{zxy} = \frac{\partial A_z}{\partial x \partial y}$ and $A_{zyy} = \frac{\partial A_z}{\partial y^2}$

Table 8 Expressions used for simulation model

2.4 Modelling of blood flow in blood circulatory system

Blood flow is the movement of blood through the vessels and is pulsatile in the large arteries, diminishing in amplitude as it approaches the capillaries. In the veins blood flow is typically non-pulsatile.

The flow of blood in arteries is the result of ventricular ejection where in the veins; it is the result of a number of factors including respiratory movement, muscle compression and the small residuum of arterial pressure.

The flow of blood in large and straight vessels, like the flow of liquid in narrow rigid tubes, is normally laminar for healthy people [38]. Laminar flow is characterized by concentric layers of blood moving in parallel down the length of a blood vessel as shown in Figure 53. The highest velocity (V_{max}) is found in the centre of the vessel. The lowest velocity ($V=0$) is found along the vessel wall. The flow profile is parabolic once laminar flow is fully developed. This usually occurs in long, straight blood vessels, under steady flow conditions.

^{§§§§} Equation 17 is different from the model used in the COMSOL reference design titled "Magnetic Drug Targeting in Cancer Therapy" in the COMSOL software. The difference had been feedback to COMSOL and engineers from COMSOL had agreed that the equation derived in this dissertation is correct.

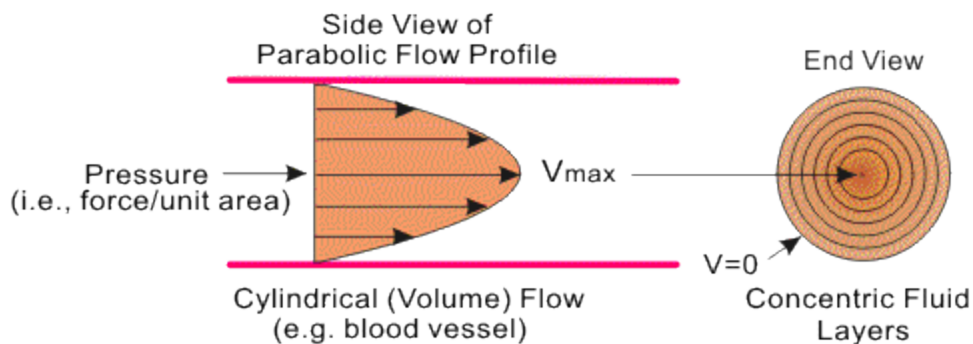


Figure 53 Illustration of laminar flow of blood in straight and large blood vessels

Turbulent blood flow is created when blood flows through a small calibre orifice and can be caused by murmurs in the heart and large arteries (Figure 54). Therefore, turbulent blood flow is usually not observed for healthy subjects.

In order to model the measurement setup for blood pulse and flow acquisition on the wrist for healthy subjects, the blood vessel on the wrist is assumed to be either the Radial or the Ulnar artery and is relatively straight as shown in Figure 55. Therefore, the blood flow can be modelled as pulsatile and having the laminar flow profile as modelled in [14].

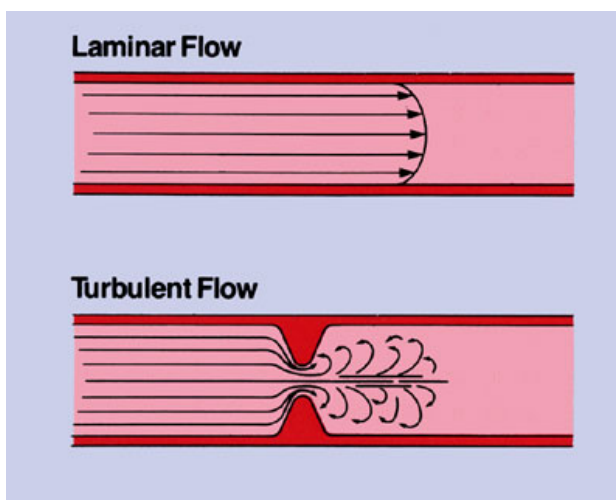


Figure 54 Diagrammatic representation of normal laminar flow in comparison with turbulent flow that results in whirls and eddies of many different velocities [39]

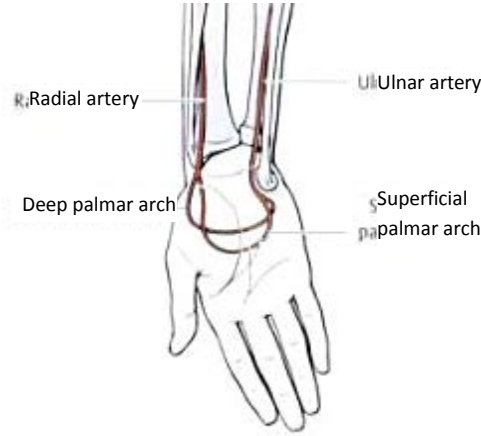


Figure 55 Illustration of arteries on the wrist

Using the assumption above, the velocity and pressure fields in the blood stream are modelled based on Navier-Stoke equations, describing the time-dependent mass and momentum balances for an incompressible flow as shown in Equation 58.

Equation 58
$$\rho \frac{\partial \mathbf{u}}{\partial t} - \nabla \cdot \boldsymbol{\eta} (\nabla \mathbf{u} + (\nabla \mathbf{u})^T) + \rho \mathbf{u} \cdot \nabla \mathbf{u} + \nabla p = \mathbf{F}$$

$$\nabla \cdot \mathbf{u} = 0$$

where η denotes the dynamic viscosity (kg/(m.s)), \mathbf{u} the blood velocity (m/s), ρ the fluid density (kg/m³), p the pressure (N/m²), and \mathbf{F} a volume force (N/m³)

The walls of the blood vessel are assumed to have no-slip condition, $\mathbf{u} = \mathbf{v} = 0$, where \mathbf{u} is the fluid velocity and \mathbf{v} is the velocity of the wall. Therefore, the outlet of the blood vessel is setup as zero pressure and the inlet boundary blood assumes a parabolic flow profile. The normal inflow velocity is accordance to $4 U_m s(1-s)$ [40], where s is a boundary segment length parameter that goes from 0 to 1 along the inlet boundary segment and U_m is the maximal flow velocity. The inflow velocity is assumed to follow a sinusoidal expression in time as described in Equation 59. In order to emulate the heartbeat, the parameters in Table 9 are used where the Equation 59 is plotted as shown in Figure 56.

Equation 59
$$U_0 = 4 \cdot U_m \cdot s \cdot (1 - s) \cdot \left\{ \sin(\omega t) + \sqrt{\sin^2(\omega t)} \right\}$$

Parameter	Value	Description
U_m	50[cm/s]	Maximum flow velocity
F	60[1/min]	Heart-beat rate
Ω	$2 \cdot \pi$ [rad]* F	Pulse angular velocity

Table 9 Summary of parameters used to model blood flow in the simulation model

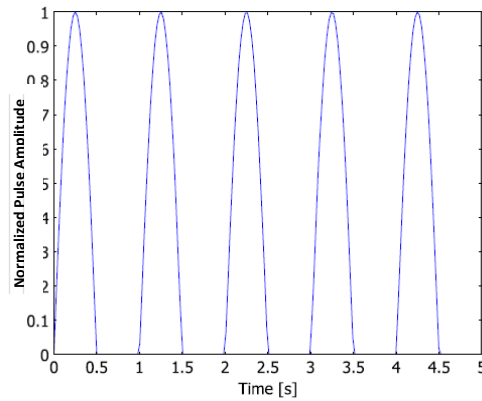


Figure 56 Input sinusoidal emulating the heart beat

2.5 Modelling of constant magnetic field

The constant magnetic field used for the MMSB experiment is based on a small cylindrical shaped permanent magnet with dimensions of 6mm diameter and 2mm height. Therefore, the key parameters used for modelling of the permanent magnet will be derived based on the need to model the magnetic field around the magnet. As such, these parameters will include the magnetic flux density, the physical dimensions and the relative permeability of the magnet.

Using the datasheet provided for the magnet, physical dimensions of the cylindrical magnet is 6mm diameter and 2mm height; the magnetic flux density is 0.2 Tesla; and the relative permeability of the magnet $\mu_{r,mag}$, is 5×10^3 as summarized in Table 10(a).

Parameter	Value	Description
mur_mag	5×10^3	Relative permeability, magnet
B_rem	0.2[T]	Remanent flux density, magnet
Height	2[mm]	Height of magnet
Diameter	6[mm]	Diameter of magnet

(a) Summary of parameters used to model magnet



Diameter, External:6mm;
Material:Neodymium Iron Boron;
Width, External:2mm;

(b) Actual magnet ***** used

Table 10 Summary of parameters used to model magnet in the simulation model

2.6 Modelling of the sensor

The magnetic sensor used for this experiment is based on the GMR sensor that is commercially available from NVE⁺⁺⁺⁺. In order to model the capability of the sensor in the transduction of magnetic to voltage, the sensor response characteristic is obtained from the sensor datasheet (as presented in Figure 12, page 23). Based on positively coupled magnetic field, multiple data points were extracted from Figure 12 and tabulated as shown in Table 11. Symmetrical modelling of the

***** <http://www.assemtech.co.uk/>

+++++ <http://www.nve.com/>

sensor characteristics in the negatively coupled magnetic field is not necessary as the MMSB model is based on stationary magnet with uni-directional application of magnetic field (i.e. north-south pole is not changed) and blood is flowing in one direction (i.e. back flow of blood is not modelled).

Applied Field (10^{-4} T)	0	5	10	15	20	25	30
Voltage (V)	0	0.075	0.15	0.23	0.275	0.285	0.29

Table 11 Table of values used to described the sensor characteristics in the simulation environment

In addition, the magnetizations of the ferromagnetic layers of the GMR sensor are reoriented relative to one another under the application of an external magnetic field. This reorientation of the magnetic moments alters both the electronic structure and the scattering of the conduction electrons in these systems, which causes the change in the resistance. As such, the GMR sensor does not generate a reactive magnetic field when an external field is applied and the final simulation environment is setup with the sensor capable of translating magnetic field into potential difference output (i.e. voltage) without the generation of reactive field.

3 System modelling of iron oxide fluidic flow in a weak magnetic field

The modelling of iron oxide fluidic flow in a weak magnetic field is based on a 2-dimensional geometry, which includes air, glass tube, permanent magnet and GMR sensor. The model, created in the multiphysics software COMSOL, will serve to present a 2-dimensional topology used in the experimental setup to demonstrate MMSB using iron oxide fluidic flow (described in Chapter 2, section 3.2).

3.1 Modelling the multiphase flow of air and iron oxide fluid

In this model, the glass tube is modelled with two inlets each with a periodic force generating a pulsatile flow of iron oxide liquid and air. At the junction where the iron oxide liquid and air mix, turbulence is generated, allowing pockets of air bubbles and iron fluid to be formed as shown in Figure 57. Using this model with sensor and coupling equations to be described in Section 2 of this chapter, the simulation was done using varying concentration of iron fluid with simulated output voltage as shown in Table 12.

Comparing simulated and measured results, the difference is found to be less than 3mV. The error can be attributed to the estimation of measured voltage, height of magnet and sensor to the glass tube and concentration of iron oxide used.

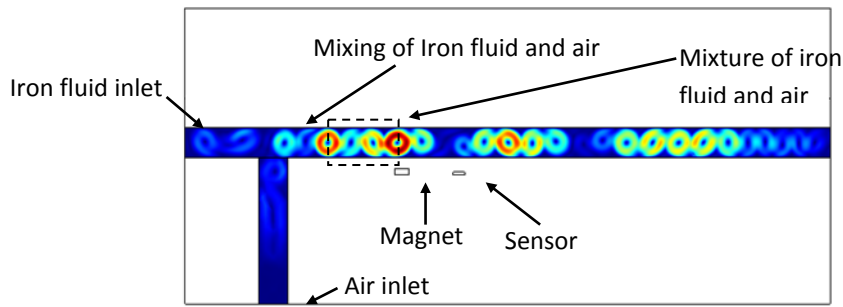


Figure 57 Two-dimensional simulation model for iron fluidic flow

Fe ₂ O ₃ concentration (g/dL)	Simulated peak-to-peak voltage (mV)	Measured peak-to-peak voltage (mV) – Estimated based on Table 3
1	11.3	13.5
3	28.1	30.0
8	40.5	43.0

Table 12 Simulation result for Fe₂O₃ fluidic flow in constant magnetic field

The successful validation of a simulation model capable of modelling magnetic disturbance was demonstrated through this simulation.

4 System modelling of MMSB

The modelling of MMSB is based on a 2-dimensional geometry to include the blood and human skin. The model will serve to present a 2-dimensional topology used in the experimental setup for MMSB signal acquisition as presented in Chapter 2, section 4. In addition, the model will also be used to verify the key parameters necessary in representing the MMSB in a simulation environment for the optimization of D_0 , the distance between the magnet and the sensor.

Using the 2-dimensional geometry illustrated in Figure 58, the domains labelled “Air” and “Tissues” are modelled as non-magnetic layers as air is known to have non-magnetic properties and the magnetic properties of tissues will be discussed in later sections. In addition, both the layers labelled “Air” and “Tissues” have large geometry and physical dimensions as compared to the other critical components in the simulation model (i.e. at least 5 times larger than the GMR sensor, permanent magnet, skin and blood vessel). Therefore, the meshing of the model for simulation will not be dependent on these parameters related to these domains.

In addition to the physical geometry, each of the critical key parameters required to model the MMSB simulation will be extracted from published information. These critical key parameters include the magnetic response of the GMR sensor; characteristics of the permanent magnet, human skin and blood; and the profile of blood flow for a healthy subject. The importance of modelling each

of these key parameters will be discussed in the following sections, where values for each parameter will be extracted or substantiated using published results.

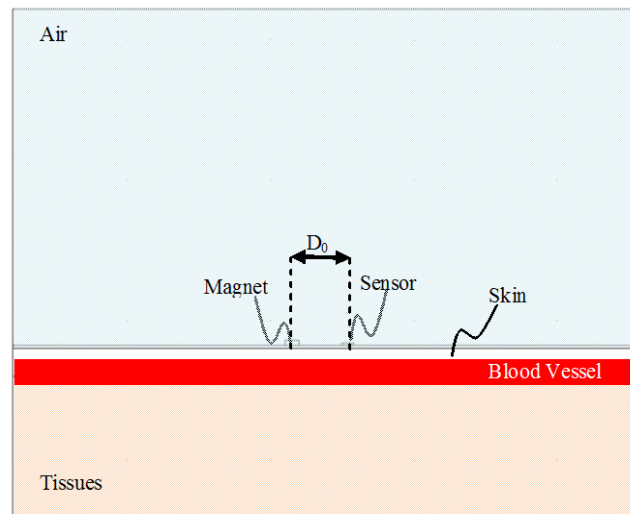


Figure 58 Illustration of the 2-dimensional simulation model geometry

4.1 Modelling the characteristics of human blood

Blood is composed of fluid plasma, formed elements, and other elements either being carried to or away from cells. Blood plasma is a liquid that serves as the extracellular environment for all the cells in the body. Total volume and concentration of plasma is important in the regulation of blood pressure. The sodium ion is the major solute in plasma, and its concentration is what determines the amount of plasma water, and thus blood volume.

The formed elements of the blood are erythrocytes, leukocytes, and platelets. Erythrocytes, or red blood cells (RBCs), are flattened discs with a depressed centre, about $2.5 \mu\text{m}$ thick and $7.5 \mu\text{m}$ wide [20]. The depressed centre provides increased surface area for the diffusion of gases. RBCs contain no nuclei or mitochondria, but they do contain the ferromagnetic particle, haemoglobin. It is the haemoglobin which binds to oxygen and allows the RBCs to transport oxygen to cells. As such, the magnetic property of blood is primarily formed by the haemoglobin, a form of iron oxides, which is present at a uniquely high concentration in the mature RBCs, while its magnetic property is affected by factors such as the state of oxygenation [21].

Therefore, blood is usually referred as the most characteristics type of biomagnetic fluid that exists in a living creature and its flow is influenced by the presence of a magnetic field. As such, in order to model blood flow in the presence of a weakly coupled magnetic field, the key magnetic parameters required are the magnetic susceptibility and permeability of blood and ferrofluidic mass content. In

addition, the conductivity, viscosity and density of the blood are also key physical parameters necessary in modelling of physical resistance to blood flow under the influence of an applied magnetic field. As such, this section will present the research done in the extraction and substantiation of these parameters through published results.

4.1.1 *Magnetic properties of blood*

Blood can be considered as a magnetic fluid because the red blood cell contains haemoglobin molecules, which is a form of iron oxides. About 30% of the red blood cell is haemoglobin. In each litre, a healthy human blood contains 3.5×10^{12} red blood cells, $4-10 \times 10^9$ white blood cells. [22]. In diseased blood the ratio of red blood cells to platelets can be much larger than 30:1.

The measure of the extent to which a material is susceptible to induce magnetization is called the magnetic susceptibility. The magnetic susceptibility is also the measure to distinguish between the types of magnetized materials. Generally speaking, a material is ferromagnetic, ferromagnetic or paramagnetic if it is attracted by a magnetic and is diamagnetic if it is repelled. Due to the difference in oxygen content in the blood, blood is reported to possess the property of diamagnetic material when oxygenated and paramagnetic when deoxygenated [23]. Measurements had been performed to estimate the magnetic susceptibility of blood and there are published reports measuring the magnetic susceptibility of blood to be 3.5×10^{-6} (i.e. weakly paramagnetic) and -6.6×10^{-7} (i.e. diamagnetic) for the venous and arterial blood respectively [14][24].

The ferrofluidic content of blood is typically referred to the amount of nanometer sized pieces of iron or various iron oxides, suspended in a liquid. In blood, the red blood cells are packed with haemoglobin and acts as the ferrofluidic content of blood. Red blood cells dominate the particular matter in blood, occupying on average about 40-45 per cent by volume of the whole blood. The percentage of red blood cells is normally given in terms of haematocrit or packed cell volume. It varies with the number and size of the red cells in the blood. The haematocrit can be defined as the volume percentage of red blood cells in the whole blood. It has been reported that the normal HCT for men 42-54% and for women, it is 38%-46% [34]. As such, the value of 0.4 is assigned to the parameter k_{ff} (or ferrofluid mass fraction in blood stream) for this model.

4.1.2 Physical properties of blood

The physical properties of blood have been studied for many years, where the Newtonian reference viscosity of blood μ_{∞} and density ρ had also reported [29] as $0.00345 \text{ N}\cdot\text{s}/\text{m}^2$ and $1050 \text{ kg}/\text{m}^3$ respectively.

In addition, blood has also been reported to exhibit considerably high static electrical conductivity, which depends on the hematocrit and the temperature. The electrical conductivity of stationary blood was measured to be 0.7 Sm^{-1} [29] while it is known that the electrical conductivity of flowing blood is always greater than that of the stationary blood. The increment for medium shear rates is about 10% and increases with the increment of the hematocrit [30]. In the study reported in [31], the electrical conductivity of blood is assumed to be temperature independent and was reported to have a value of 0.8 Sm^{-1} . As such, the electrical conductivity of blood is assumed to be 0.8 Sm^{-1} in the proposed MMSB model.

4.1.3 Summary

Using published reports on the properties of blood, the model to simulate the phenomena of MMSB will be based on these reported values as summarized in Table 13.

Parameter	Value	Description
chi_ff	-0.667×10^{-7}	Magnetic susceptibility of ferrofluid (blood)
mur_ff	$1 + \text{chi_ff}$	Relative permeability of ferrofluid (blood)
k_ff	0.4	Ferrofluid mass fraction in blood stream
Rho	$1050 [\text{kg}/\text{m}^3]$	Density, blood
μ_{∞}	$0.00345 [\text{N}\cdot\text{s}/\text{m}^2]$	Newtonian reference viscosity, blood
Σ	$0.8 [\text{Sm}^{-1}]$	Electrical conductivity of blood

Table 13 Summary of parameters used to model blood in the simulation model

4.2 Modelling of the human skin

In order to model the human skin for MMSB phenomena, the skin thickness and its magnetic properties are important considerations. This section will present the findings of data related to skin thickness and its magnetic properties in order to support the modelling of MMSB phenomena in a multi-physics simulation environment.

4.2.1 Skin thickness on the wrist

Standard skin thickness measurements are meant to describe the thickness of the epidermis and dermis. These layers are firmly adhered between the papillary layers of the dermis and stratum germinativum of the epidermis. The thickness varies from 0.5 – 4mm. Skin is generally divided in terms of thickness into thin skin and thick skin. Thick skin is found on the palms of the hand and soles of the feet (4mm) with thin skin covering the remainder of the body, thinnest skin is eyelid 0.5mm. The variations in skin thickness within the body are due almost entirely to the variation in thickness of the dermis with the dermis of the extensor surfaces (back of the forearm – posterior surface) generally being thicker than the flexor surfaces (front of the forearm anterior surface).

According to several authors (example: Becker 1939; Southwood 1955), the thickness of the skin on the palmar forearm varies from 1.2 – 1.7mm [35]. However, the skin on the wrist should be slightly thinner as we move away from the palm towards the arm. Therefore, in order to account for the deviation of the skin thickness reported on the palmar forearm and the wrist, the thickness of the skin layer used for modelling of MMSB will be based on 1mm.

4.2.2 Magnetic properties of the skin

It has been reported [36],[37] for DC magnetic (H) field, the magnetic permeability μ of most biological materials is practically equal to the magnetic permeability μ_0 of free space. However, time-varying H field may interact with living organisms and slightly modify this property.

Since MMSB signal variation is primarily based on the position of a permanent magnet with close proximity to the blood vessel and GMR sensor, the applied field is a stationary H field and the human skin will be modelled with the magnetic permeability of free space, μ_0 (i.e. $4\pi \times 10^{-7}$ H/m).

5 Coupling between ferro-fluidic volume force equation with applied magnetic field

Magnetic transport of red blood cell in the vascular system is governed by several factors including (a) the magnetic force, (b) viscous drag, (c) red blood cells interactions, (d) inertia, (e) buoyancy, (f) gravity, (g) thermal kinetics (Brownian motion), (h) red blood cells / bio-fluid interactions (perturbations to the flow field), and (i) inter-red blood cell effects such as magnetic dipole interactions. In this section, it is assumed that the dominant factors are magnetic and viscous forces, which will be taken into account to explain the coupling between ferro-fluidic volume force and magnetization force in COMSOL simulation.

The induced magnetization of a ferro-fluid in the x and y axes had been derived in Equation 56. Using the magnetization M_x as an example, it can be observed that a red blood cell moving in the x-axis will induce a magnetization of $M_x = \frac{\chi}{\mu_0} \frac{\partial A_z}{\partial y}$. As such, the magnetic field generated by the red blood cell will produce a magnetic field that is dependent on the magnetic field A_z , which is applied perpendicularly to the direction of blood flow using the permanent magnet.

To model the movement of the red blood cell under the ferro-fluidic volume force, Equation 57 is derived. Each of these ferromagnetic red blood cell will experience the ferro-fluidic volume force in the x-axis $F_x = \frac{\chi}{\mu_r \mu_0^2} \left(\frac{\partial A_z}{\partial x} \frac{\partial^2 A_z}{\partial y^2} + \frac{\partial A_z}{\partial y} \frac{\partial^2 A_z}{\partial y \partial x} \right)$, which is a function of the applied magnetic field and the displacement in the blood vessel.

The rate of displacement for the red blood cell into the blood vessel is modelled by Equation 59, which defines the inlet pressure. The distribution of the red blood cell in the blood vessel will follow the Navier-Stroke Equation 58, which models the velocity and pressure fields in the blood stream under no-slip and non-turbulence condition.

Coupling Equation 59, Equation 58, Equation 57 and Equation 56 in COMSOL for simulation, the magnetization is solved for each time step, where the ferromagnetic red blood cell is displaced into the blood vessel, generating a measurable magnetic disturbance at the sensor location through the magnetization vector, as a result of the applied perpendicular magnetic field by the permanent magnet. The simulated result from COMSOL, based on these coupling, will be described in the section 6 of this chapter.

6 Numerical solutions of 2D model for MMSB simulations

In order to simulate the model created in COMSOL, a numerical solution is required where non-linear meshing of the model is necessary to provide the resolution at the location near the sensor and magnetic while most of the other parts of the model can be meshed using larger meshing dimensions. This can be achieved using the triangular meshing element, or the equivalent of tetrahedral edge elements for 3-D models.

The triangular meshing element is selected for its ease of convergence [43] and ability to achieve non-linear meshing of the gaseous (air), solid (skin and tissues) and liquid (blood) domains with magnetic coupling. In addition, it has been reported in [44], that the numerical dispersion error is very low for meshes composed of well-shaped tetrahedral edge elements in electromagnetic modelling using the finite-element method. Therefore, using the non-linear triangular meshing element, the 2-D model of Figure 58 is meshed as shown in Figure 59 and Figure 60.

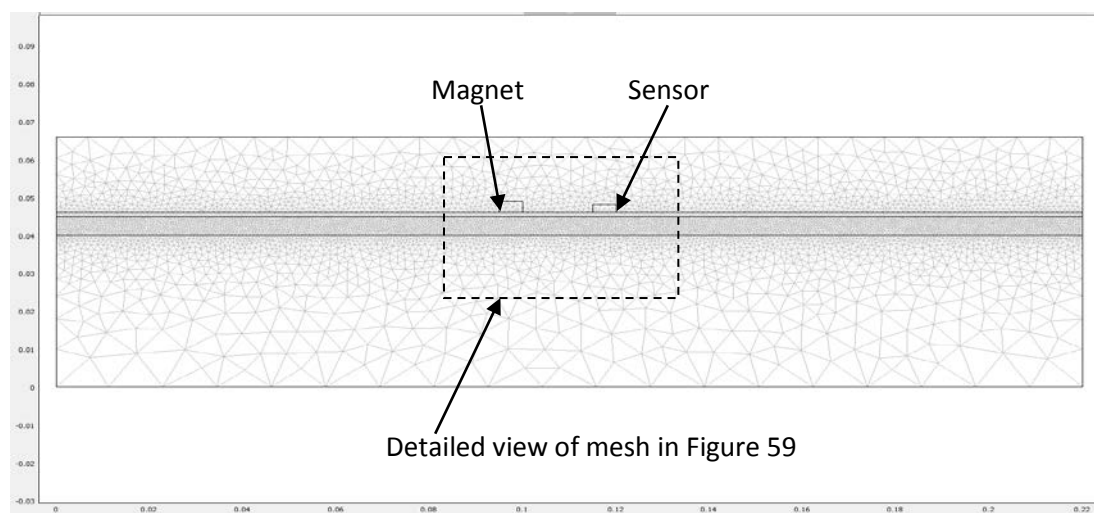


Figure 59 Meshing results for 2D MMSB model using multi-physics simulation software COMSOL

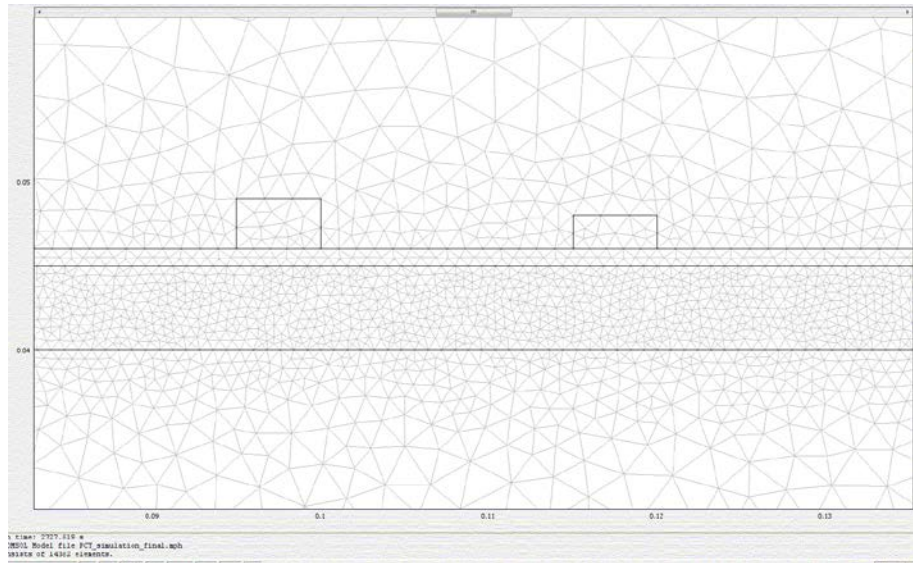


Figure 60 Detailed view of meshing results near sensor and magnet using multi-physics simulation software COMSOL

Having the 2D model created and meshed in COMSOL using the parameters described in this chapter, a transient (quasi-static) simulation is executed for a period of 2 seconds with time step of 0.05 seconds. The waveform obtained is shown in Figure 61, includes the amplifier gain of 100 so as to produce comparable results to the physical experimental setup used in Figure 27.

From the waveform obtained in Figure 61, the 2D model is verified to be capable of modelling MMSB as the output signal (Figure 61) follows similar waveform characteristics as the input stimulus (Equation 59).

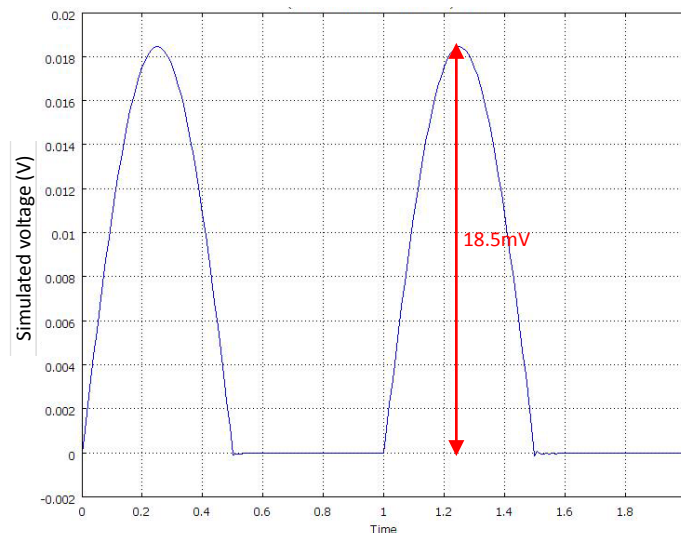


Figure 61 Simulation results for 2 seconds based on a distance of 20mm between magnet and sensor

To check that the 2D model created in COMSOL is giving similar results as the physical experiment, the output voltage and total pulse duration of the measured MMSB waveform is analyzed as shown

in Figure 62. Using the ventricular response of the experimentally measured waveform (Figure 62) as the reference, it can be observed that the ventricular response has a peak voltage of 19mV (approximately) and a total pulse duration of 0.5s. This is similar to the simulated waveform from COMSOL as shown in Figure 61 where the peak voltage simulated is 18.5mV with a waveform periodicity of 1 second and total pulse duration of 0.5s.

Therefore, comparing the results in Figure 61 with Figure 62, it can be concluded that the 2D model created in COMSOL is capable of modelling the MMSB phenomena on human wrist (skin) based on published information available on the parameters and equations listed in Section 2 and 3 of Chapter 2 respectively.

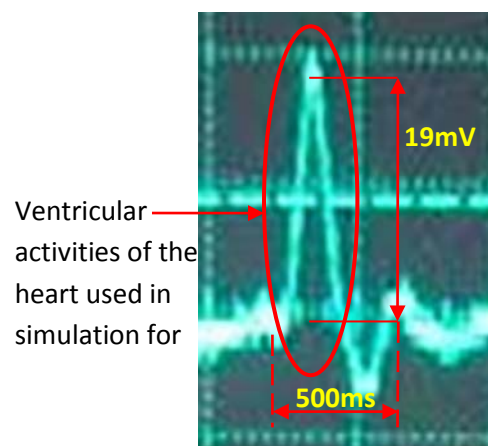


Figure 62 Waveform captured from the sensor output using oscilloscope for comparison with simulation

With the 2-D model verified to be capable of simulating the MMSB phenomena on human subjects, the distance between the sensor and the magnet (Figure 58), D_0 , is varied. The simulated output voltages of the sensor (after amplification) are recorded and plotted as voltage (normalized) as shown in Figure 63. From Figure 63, it can be observed that the optimal distance, D_0 , between the magnet and sensor is approximately 2cm, where the output voltage is highest. This will be verified in the next section where a series of experiments were conducted on human subjects to validate the simulation result.

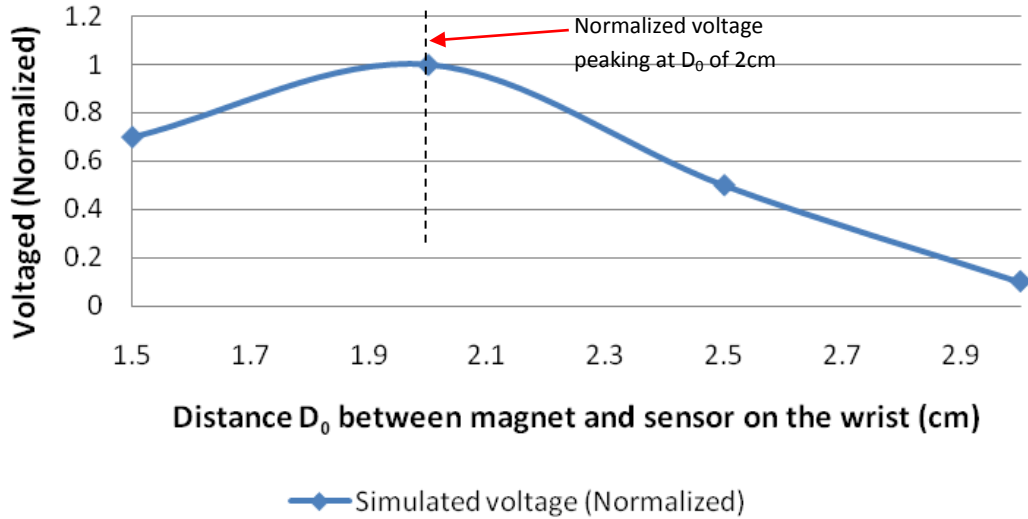


Figure 63 Plot of simulation results with varying distance between sensor and magnet

7 Blood pulse measurements using MMSB for 20 subjects

Using the measurement setup described in Chapter 2, Figure 26(a) and Figure 27, a trial was conducted on 20 subjects aged 20 – 24, to understand the statistical difference of the measured data on different human subjects. The distance between the magnet and the sensor was chosen as 1.5cm, 2cm, 2.5cm and 3cm. This is because the magnet has a diameter of 6mm. In order for repeatability of measurement results, it was found that a small displacement (i.e. < 5mm) of the magnet away from the sensor tends to result in the sensor slipping and falling off the skin. In addition, for some cases, measurements on subjects with the use of sticky tapes was found to have skin allergies and results for repeated measurements tend to vary for very in-consistent. As such, in this experiment, each displacement of the magnet from away from the sensor is set as 5mm as this is close to the diameter of the magnet and repeatability of measurement results is possible. In addition, with the displacement selected at each step of 5mm, positioning of the magnet on the wrist is also easier, where slipping of magnet off the wrist can be better managed without the use of the sticky tape, resulting in the possibility of minimising discomfort and potential allergies to the test subject.

The measurement results are plotted in both radar chart (Figure 64(a)) and column chart (Figure 64(b)) to illustrate the probability of data peaking at a particular distance and also the trend of the data measured, respectively. Using the radar chart (Figure 64(a)), it can be observed that the measurement results have a relatively high probability of peaking at a distance of 2cm between the magnet and sensor as compared to other distances. This can be further analyzed in (Figure 64(b)) where the addition of the trend-line in the column plot clearly identifies the maximal of the trend-

line at an optimal distance of 2cm between the magnet and sensor to produce maximal MMSB signal.

With the statistical analysis completed based on the trial conducted, the optimal distance of 2cm between the magnet and sensor to produce maximal MMSB signal will be confirmed with a comparison with the 2D simulation model developed in COMSOL.

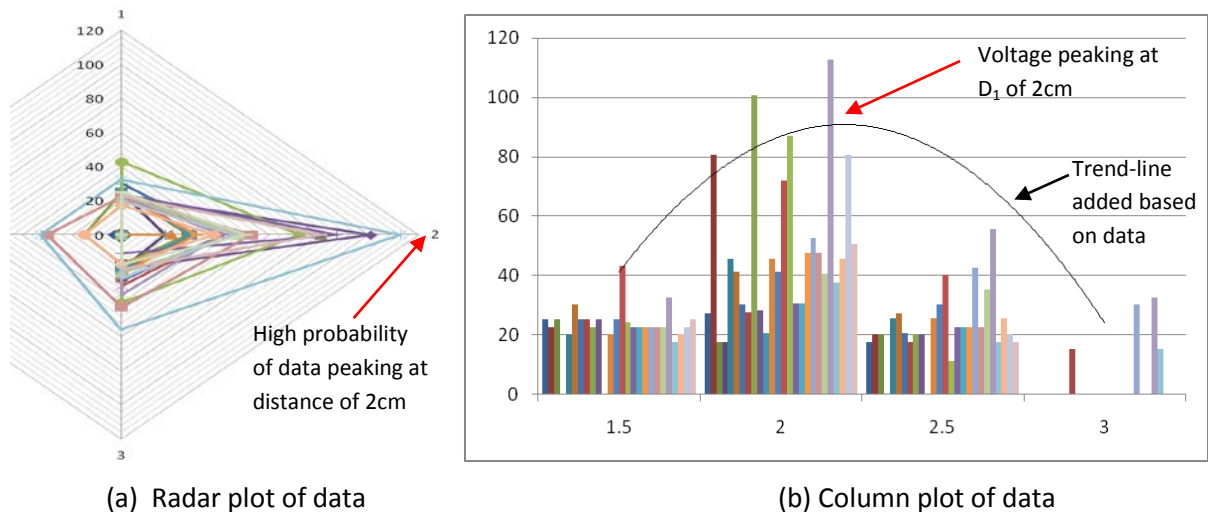


Figure 64 Plots of distance vs measured voltage based on measured data from 20 subjects

8 Comparison of measured and simulated results

In order to compare the measured and simulated results, the voltages obtained, for both measured and simulated results, at each of the distance 1.5cm, 2cm, 2.5cm and 3cm are averaged and normalized to the maximum value as tabulated in Table 14. The normalized results are plotted as shown in Figure 65.

Distance (cm)	Simulated voltage (Normalized)	Measured voltage (Normalized)
1.5	0.701389	0.465181
2	1	1
2.5	0.500102	0.475925
3	0.094859	0.073617

Table 14 Tabulated results for simulated and measured data

Using Figure 65, it can be observed that the simulated and measured data both peak at an approximate distance of 2cm between the sensor and the magnet. In addition, as the distance between the sensor and magnet increases, both the simulated and measured voltages decline at a similar rate showing good correlation of the two results.

However, the measured and simulated results at the distance of 1.5 cm between the sensor and magnet are not the same. This can be attributed to errors in the measurement setup where at close proximity between the sensor and magnet, the accuracy in maintaining the smaller distance is lower. Nevertheless, the results at 1.5cm are still lower than the results at 2cm confirming the optimal distance between the sensor and magnet is 2cm based on the measurement configurations used.

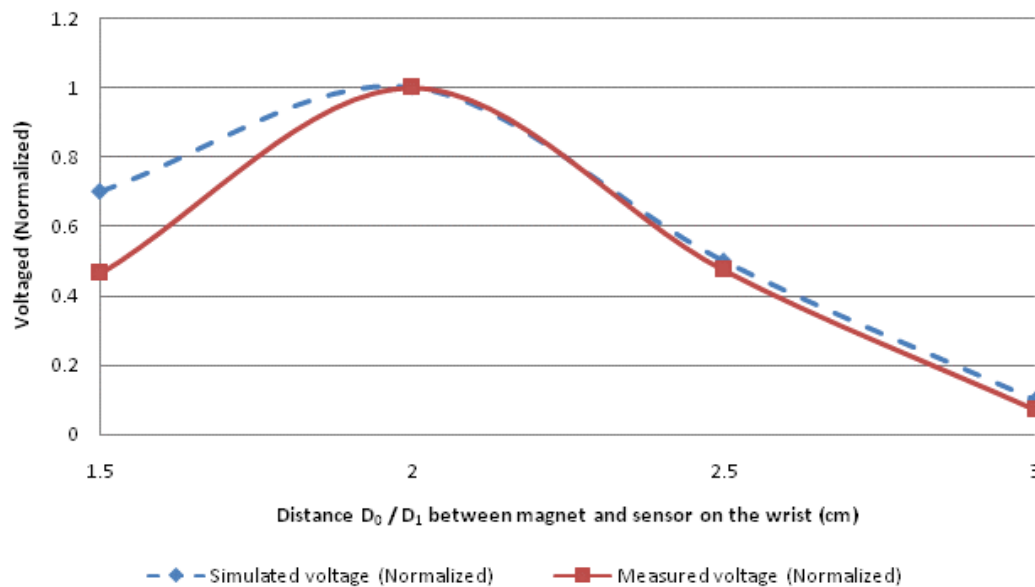


Figure 65 Comparison of simulated and measured voltage (normalized) with varying distance between sensor and magnet

9 Conclusions

The simulation model presented in this chapter is based on the MMSB phenomena proposed in this dissertation. The simulation model is found to be highly correlated with experimental results and it can be concluded that the simulation model has successfully modelled the MMSB phenomena. Such a model will allow the sensitivity of the MMSB method to be assessed for different human subjects and different parts of the body. In addition, the model can also be used to achieve design optimization of the distance D_0 , between the sensor and the magnet, so as to achieve robustness in the physical implementation of MMSB as a wearable device for pulse rate measurements.

With the completion of MMSB modelling in a simulation environment, the next chapter will present the findings on the applications of MMSB for blood flow. In addition, the blood flow result is verified using the post-occlusive reactive hyperaemia (PORH) test with an instrument capable of Laser Doppler Flowmetry.

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Chapter 4 Measurement of blood flow using MMSB

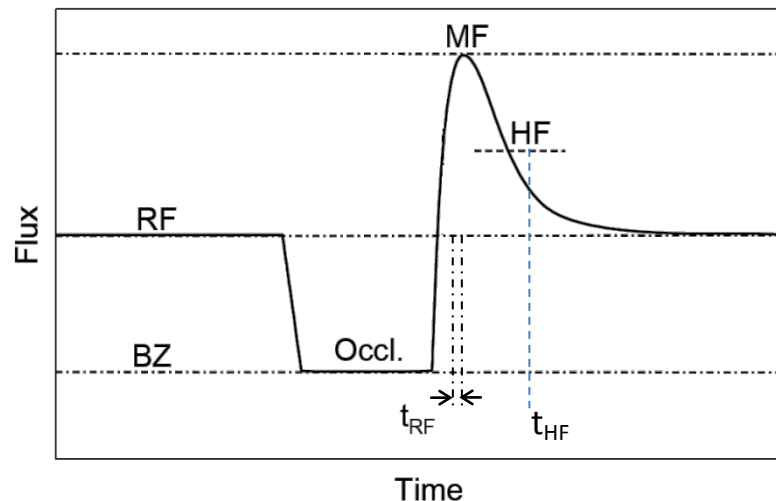
1 Introduction

Blood flow refers to the flow of blood in the cardiovascular system and is typically measured by observing blood flowing through a section of the blood circulatory system. It is a physiological sign that is dependent on the heart pushing blood into the arteries and the resistance present in the arteries, which will expand and contract allowing blood to flow.

Blood flowing in the skin is one of such physiological signs, which varies from individual to individual, is site dependent and is affected by a combination of systemic, local and disease-related factors.

Endothelial function may also be a determinant of skin perfusion. If a decrease of blood flow occurs, the increase of waste products or decrease of oxygen levels will induce vasodilatation that will enhance blood flow until the normal situation is reached again. In healthy subjects, their regulation system is able to increase the blood perfusion at least ten-fold compared to typical resting flow conditions [1]. However, in the case of patients with Peripheral Arterial Obstructive Disease (PAOD), the presence of stenosis in proximal arteries will decrease the available pressure at the level of the arterioles. In this type of cases, distal vasodilatation is an appropriate reaction to prevent an increased concentration of waste products (i.e. carbon dioxide) and to supply enough oxygen in order to maintain the necessary metabolic condition.

In order to assess the microvascular function of human, the post-occlusive reactive hyperaemia (PORH) test using laser Doppler perfusion monitoring has been proposed in [2]. The local blood perfusion, e.g. at a distal extremity, is measured before, during and after performing arterial occlusion to record the response upon releasing the occlusion. The measurement is typically performed by the means of a Laser Doppler Perfusion Monitor (LDPM) and a typical signal output is illustrated in Figure 66 [3].



BZ = biological zero, Occl. = occlusion period, RF = resting flux, MF = maximum flux, HF = half flux decrease between maximum and resting flux, t_{RF} = time of cross point with RF, t_{HF} = time to half return to resting flux after maximum, measured from time of cuff release

Figure 66 Typical PORH response of a healthy subject [1]

Upon occlusion, a PORH tracing shows that the blood perfusion drops from its resting flux values to the biological zero (BZ) level [4]. After releasing the cuff the blood perfusion will return to the resting flux (RF) value with an overshoot. The magnitude and time regime of that overshoot are clinically relevant diagnostic variables [5][6].

In normal (healthy) cases, immediately after cuff release the flux will quickly rise to a maximum (MF) followed by a slower decrease to initial (resting flux) values as shown in Figure 67(a) [7]. The reason for the increased flux after cuff release is mostly the vasodilatation due to the ischemia produced by the occlusion. Both endothelium-dependent and –independent effects might modulate the degree of vasodilatation. In the presence of proximal stenosis or obstruction (atherosclerosis), the peak perfusion value that occurs after occlusion will be relatively lower. In case of critical ischemia, the maximum vasodilatation has already been reached in the resting period, before occlusion, and the PORH response will not show a higher response than that resting flux level. Also the increase of flux after occlusion will be smaller because of the presence of stenosis in the larger arteries, which will inhibit a rapid refilling of the arterial segments.

The rate of flux increase (t_{RF}) just after cuff release is another important parameter, which is much lower in the case of PAOD. After the occurrence of the maximum flow, the oxygenation of the tissue normalizes, and the need for the hyperaemic response vanishes. At that time, the vasodilatation will decrease and be stabilised back to resting flow (RF) as illustrated in Figure 67(b) [7]. Typical result for healthy subjects has been reported [2], at the 95% confidence interval for the mean of t_{RF} , was found

to be approximately a 10th order of magnitude between healthy and PAOD patients. This can also be observed in the PORH waveform in Figure 67.

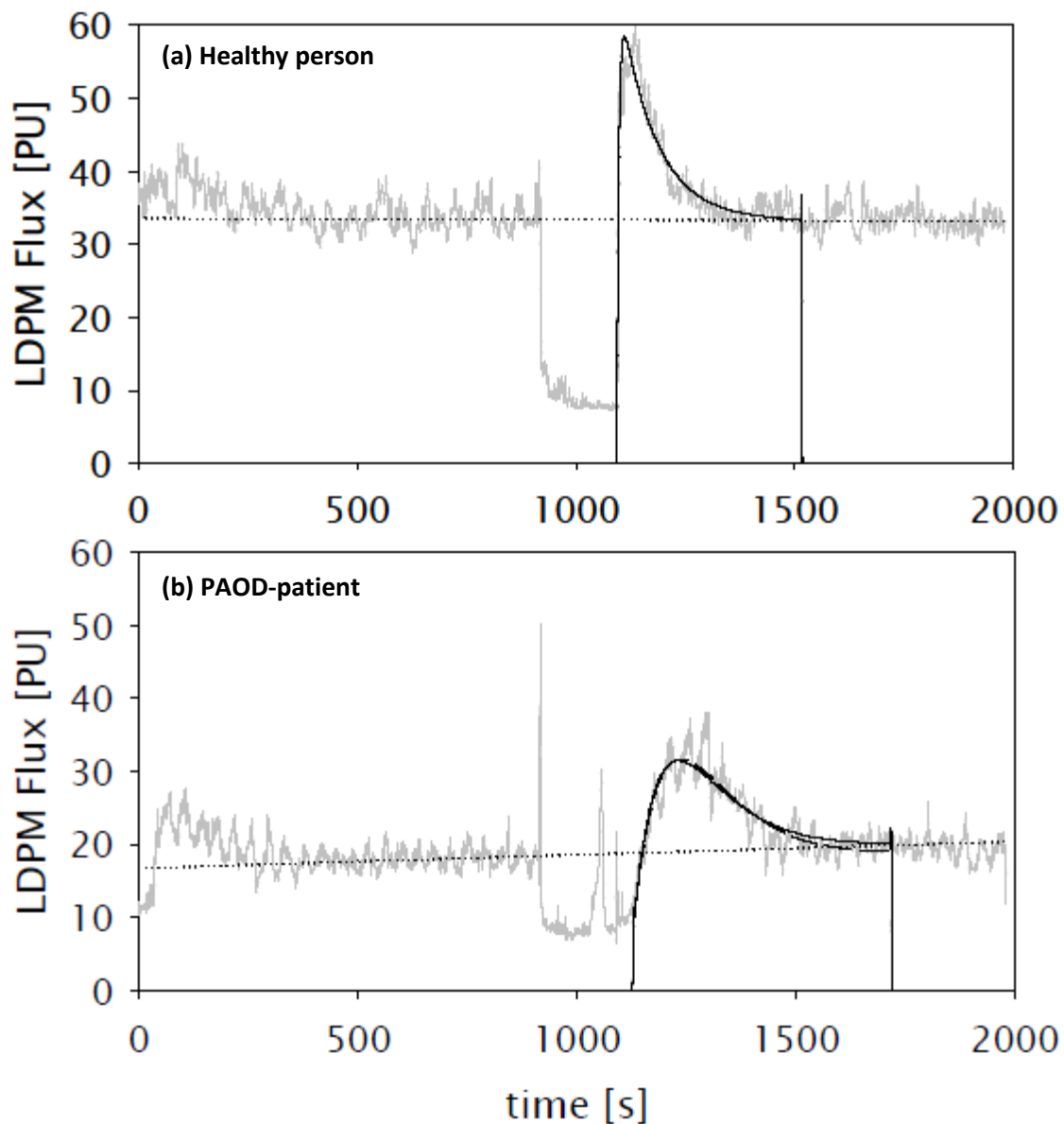


Figure 67 Analysis of typical PORH-recordings, from a healthy person (a) and a PAOD-patient (b) [7]

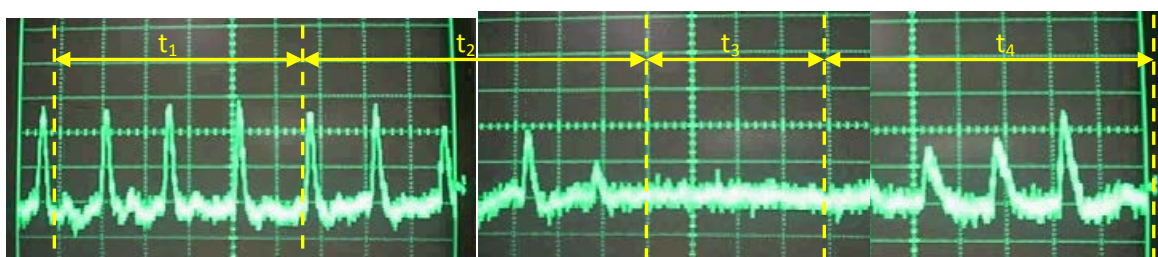
Current methods of PORH measurements are optical based and requires direct contact with the human skin with reasonably good optical properties (i.e. good optical reflectance from the skin). As such, performance of these devices tend to vary with time and are subjective to the presence of human body fluids (e.g. blood) and environmental contaminants (e.g. mud, water, etc). This chapter proposes the application of the novel method, MMSB, to acquire blood flow non-invasively, using the magnetic disturbance created by blood flowing through the localized magnetic field. Using the MMSB phenomena, blood flow in the skin can be detected using a portable configuration capable of

measurements through non-magnetic material such as fabric. This will be able to support research in autoregulatory mechanisms where studies on the affects of blood flow during and following pressure assault are useful for pressure ulcer aetiology.

2 Experimental setup for blood flow measurements using MMSB

Based on the principles of MMSB, the approach to interpret the amplitude of the output waveform from the GMR sensor (as shown in Figure 29, Chapter 2) is to assume a functional relationship exists between the amplitude and the intensity of blood flow. This is because an increase in blood flow intensity will create a larger disturbance to the uniform magnetic field which can be picked-up by the GMR sensor as the output voltage amplitude. However, this approach is based on the assumption that the sensor is magnetically biased to operate along the linear region as discussed in Figure 12, Chapter 1.

In order to verify the approach in interpreting the amplitude of the output waveform from the GMR sensor, an experiment was conducted using an inflatable cuff placed on the forearm. The experimental setup in Figure 25, Chapter 2, is designed to verify that the voltage of output waveform for the MMSB has a functional relationship to the difference in blood flow. The GMR sensor and magnet was placed on the wrist as the pulsatile effect of the arteries will allow blood flow to be measured effectively [8]. The cuff is automatically pressurized and released using the digital monitor for blood pressure. During this process of pressurization and releasing of the pressure in the cuff, the output voltage of the GMR sensor is measured as shown in Figure 68.



- t₁: Period of MMSB measurements without cuff activation
- t₂: Period of MMSB measurements with cuff pressure increasing
- t₃: Period of MMSB measurements with cuff pressure holding
- t₄: Period of MMSB measurements with cuff pressure releasing

Figure 68 Signal captured using MMSB with blood flow restriction created with the use of the digital monitor for blood pressure measurements

Based on the waveform obtained in Figure 68, it can be observed that the output voltage is changing with respect to the increase in pressure for the cuff on the forearm. The waveform can be

segmented into four sections, namely t_1 , t_2 , t_3 and t_4 each signifying a section of the waveform critical in the analysis of the use of MMSB for blood flow measurements.

During the period of t_1 , the amplitude of the voltages of each pulse is similar signifying the consistency of blood flow with no changes of the pressure on the cuff.

However, in the period marked as t_2 , the pressure in the cuff starts to increase. At the same time, it can be observed that the pulse amplitude also starts to decrease. This continues till the pressure in the cuff is sufficient to limit the flow of blood in the arm resulting in a non-measurable output voltage by the GMR sensor. With this decrease in amplitude due to the increase in pressure in the cuff, it can be concluded that the decrease in blood flow due to the increase in pressure in the cuff has resulted in the weakening of the MMSB phenomena resulting in the decrease in the pulse amplitude measured.

With the cuff pressure increased to its maximum in the period t_3 , the blood flow will reduce to minimum resulting in minimal or non-detectable MMSB. This is another conclusive observation that the MMSB phenomenon is correlated to the blood flow.

Finally, during the period of t_4 , the pressure on the cuff starts to release and blood starts to flow again into the forearm and the wrist. With the increase of blood flow due to the release of the pressure in the cuff, the output voltage for the pulse detected using the MMSB becomes observable again and is increasing in amplitude with respect to the rate of the pressure release in the cuff.

From this experiment, it can be concluded that the amplitude of the voltage measured from the GMR sensor using the MMSB phenomena is correlated to the amount of blood flowing in the arteries in an inversely proportional manner. In order to verify this inverse proportionality, the following section will enhance the model created in Chapter 3 by including an exponential decreasing component in the blood flow equation, where the voltage output of the GMR sensor will be plotted with respect to time.

Upon verification of the correlation between MMSB and blood flow analysis, the rest of Chapter 4 will apply the MMSB and Laser Doppler method for PORH measurements.

3 Simulation of blood flow using MMSB

Using the simulation model created in Chapter 3, the normal inflow velocity model for blood was described by Equation 59 in Chapter 3. This equation is modified in this section to simulate the

change in blood flow due to the application of occlusions through the pressured cuff. The increase in applied pressure will reduce the blood flow in the lower arm. This phenomenon can be modelled as a decrease in amplitude of the normal inflow velocity of the blood over time.

In order to describe the reduction of blood flow over time, an exponential decay term is applied to Equation 59 as shown in Equation 60. The exponential decay term, $e^{-0.1t}$, in Equation 60 models the rate of decay based on the measured MMSB amplitude in Figure 68. The measured MMSB peak amplitude was observed to decrease at an approximate rate of 10% between each MMSB peak. Therefore, the inclusion of the exponential decay term will simulate the effect of the pressurized cuff inflating and thus, reduction in blood flow.

$$\text{Equation 60} \quad U_0 = 4. e^{-0.1t} \left(U_m \cdot s. (1 - s). \left\{ \sin(\omega t) + \sqrt{\sin^2(\omega t)} \right\} \right)$$

Using Equation 60 as the enhanced blood flow model of Equation 59, the MMSB simulation was executed for a period of 10 seconds. In this simulation, the output voltage of the GMR sensor was recorded with respect to time and shown in Figure 69.

From Figure 69, it can be observed that the simulated output voltage of the GMR sensor decreases in amplitude with time at a proportion of 10% from the previous peak. This is similar to the measured MMSB waveform in Figure 68 demonstrating the proportional reduction of magnetic disturbance with respect to an increase in the applied pressure. As such, it can be concluded that the enhanced blood flow model is capable of simulating the reduction in blood flow due to an externally applied pressure (e.g. pressurized cuff).

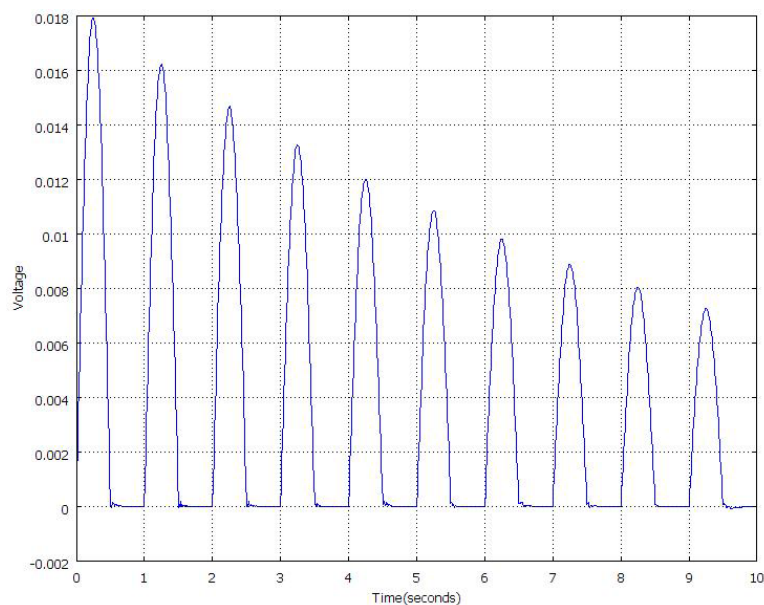


Figure 69 Simulated MMSB output voltage using modified equation for inflow velocity of blood

4 PORH measurements setup using laser Doppler medical instrument

The experiment setup for acquisition of blood flow in the skin using the PORH test is based on the Laser Doppler Flowmetry instrument DRT4, from Moor Instruments, designed specifically for advanced clinical and research applications. Laser Doppler Flowmetry (LDF) works by illuminating the tissue under observation with low power laser light from a probe containing optical fiber light guides. Laser light from one fiber is scattered within the tissue and some is scattered back to the probe. Another optical fiber collects the backscattered light from the tissue and returns it to the monitor. Most of the light is scattered by tissue that is not moving but a small percentage of the returned light is scattered by moving red blood cells. The light returned to the monitor undergoes signal processing whereby the emitted and returned signals are compared to extract the Doppler shift related to moving red blood cells. The extracted Doppler shift is processed by the instrument DRT4, which has full computer connectivity to collect and store the data from automated measurements involving pressure cuff provocations for PORH.

A typical measurement setup is shown in Figure 70 where an inflatable cuff is wrapped around the forearm above the elbow. The laser probe is placed below the inflatable cuff to acquire blood perfusion data. An occlusion is applied on the cuff via the DRT4 instrument where the auxiliary artery was compressed automatically for 20 seconds and then released. The changes in flow flux are displayed in real time to ensure complete occlusion before the inflated cuff is released.

Concurrently, data is acquired on a computer to record the flow flux measured by the laser Doppler instrument before the occlusion till 1 minute after occlusion as shown in Figure 71 where the resting flux (RF) can be clearly identified. At time t_b , the cuff occlusions occur and a peak signal is induced in the measured results due to the motion artefacts created. This is followed by the occlusion period itself and the maximal flux (MF) peak as marked in Figure 71.

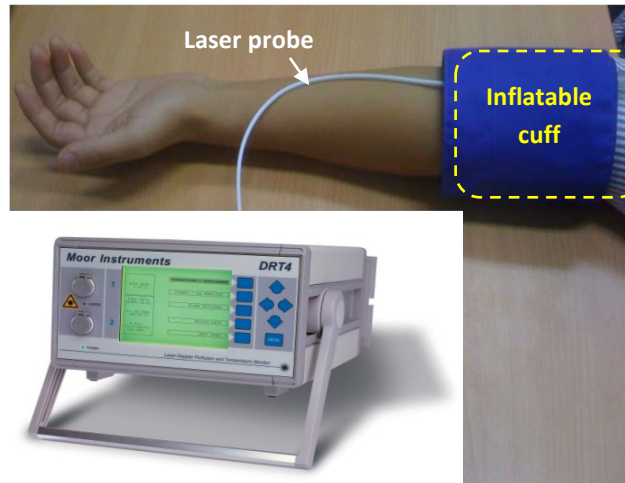


Figure 70 Illustrations of typical measurement setup for PORH using DRT4, Moor Instrument

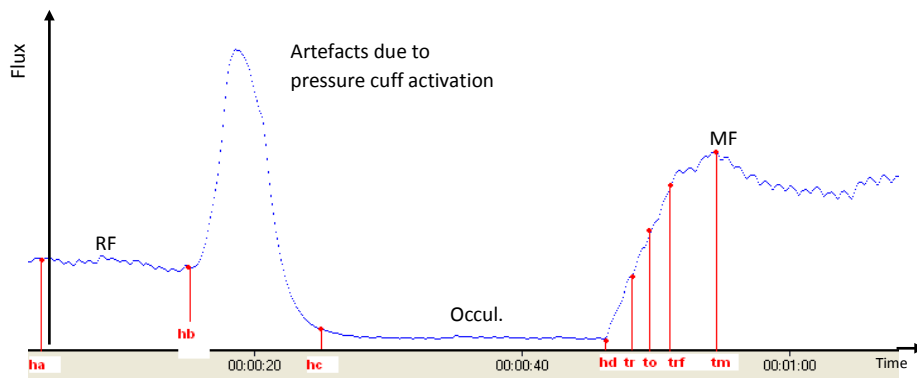


Figure 71 Typical results obtained using the laser Doppler Flowmetry method for PORH measurements

5 PORH measurements setup using MMSB acquired on the wrist

The alternative and new experiment for acquisition of blood flow in the skin utilizes the concept of placing a magnetic field in the vicinity of the major artery where the blood flowing through the magnetic field will disturb the magnetic field, thus creating a magnetic disturbance.

Using the optimal distance of 2cm obtained in Chapter 3 for the distance between the magnet and sensor, the set-up for the PORH measurement is shown in Figure 72. The sensor output is sufficiently amplified and then connected to an oscilloscope. Data is acquired using the National Instrument data acquisition card (NI USB-6008) with 12 bit Analog-to-Digital resolution at a sampling rate of 1 kHz using Figure 28 in Chapter 2.

The occlusion is simulated using the pressure cuff from the Moor Instrument DRT4 and concurrent measurements are done, in-parallel, using MMSB setup and laser Doppler probe as shown in Figure 72. The waveform obtained using the MMSB sensor is shown in Figure 73 where the pulse amplitude

is observed to be closely correlated to the rate of pressure increase and decrease by the inflatable cuff. In addition, it can be observed from Figure 73 that there exist well identified RF, occlusions and MF zone. However, with the cuff placed further away from the sensor as compared to the laser probe, the motion artifacts due to pressure cuff activation, is not observable as compared to results from the laser Doppler method in Figure 71.

In order to have a better understanding on the flow of blood during the pressurization and depressurization of the cuff, the waveform in Figure 73 is used where the measured result is zoomed in for the time period from the moment before pressurization of the cuff to the time when maximum pressure is exerted in the cuff resulting in occlusions. The result obtained is shown in Figure 74 where the increase in pressure in the cuff resulted in an observable decrease in the amplitude of the blood pulse. In addition, from Figure 74, it can be observed that the amplitude of the blood pulse decreases due to pressurization of the cuff until it is not observable before occlusions.

Similarly, using Figure 73, the measured result is zoomed in for the time period from the moment before depressurization of the cuff to the time when maximum flow (MF) is observed. The result obtained is shown in Figure 75 where the decrease in pressure in the cuff resulted in a observable increase in blood flow. In addition, from Figure 75, the amplitude of the blood pulse becomes observable as blood flow increases due to the depressurization of the cuff.

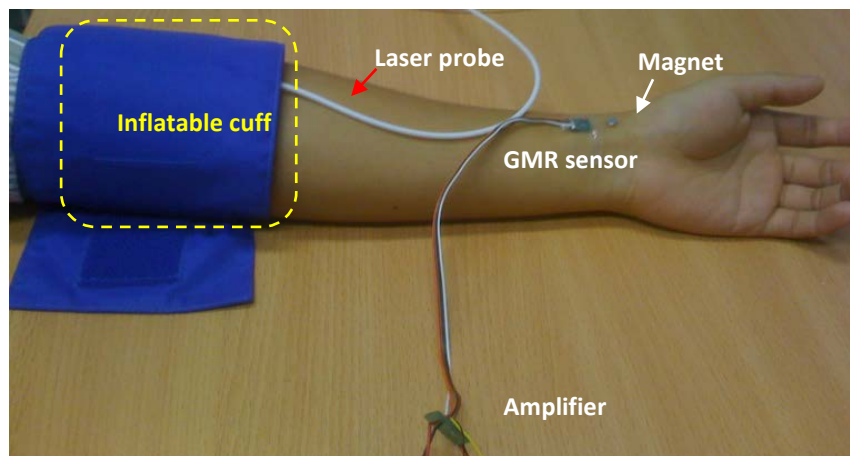


Figure 72 Illustration of blood flow acquisition on the wrist

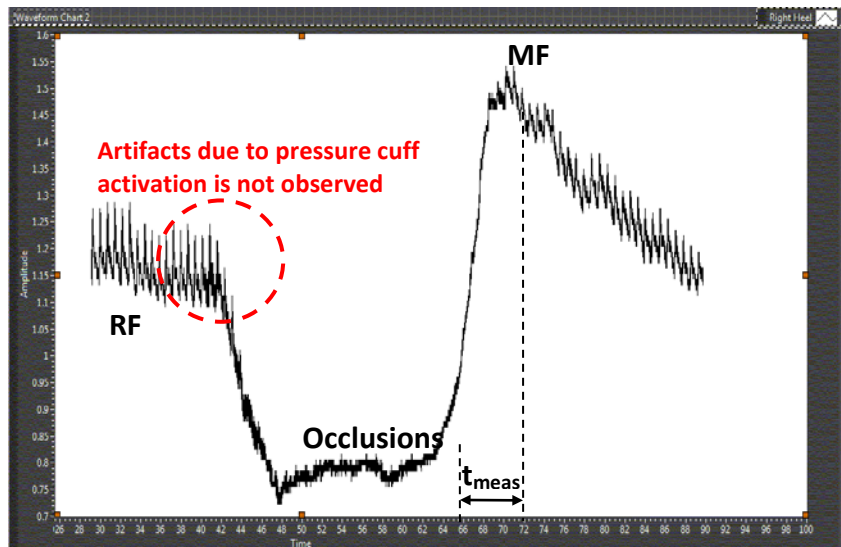


Figure 73 Typical results obtained using the MMSB for PORH measurements

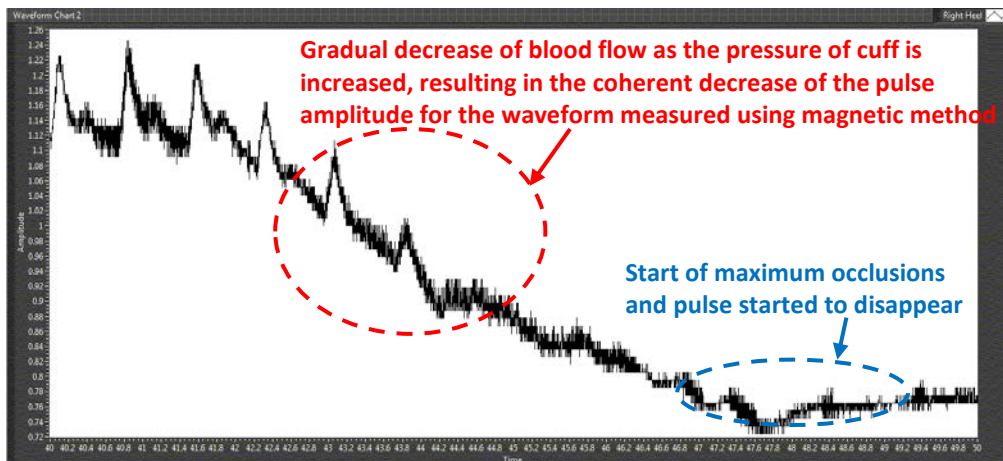


Figure 74 Waveform obtained using MMSB before occlusion

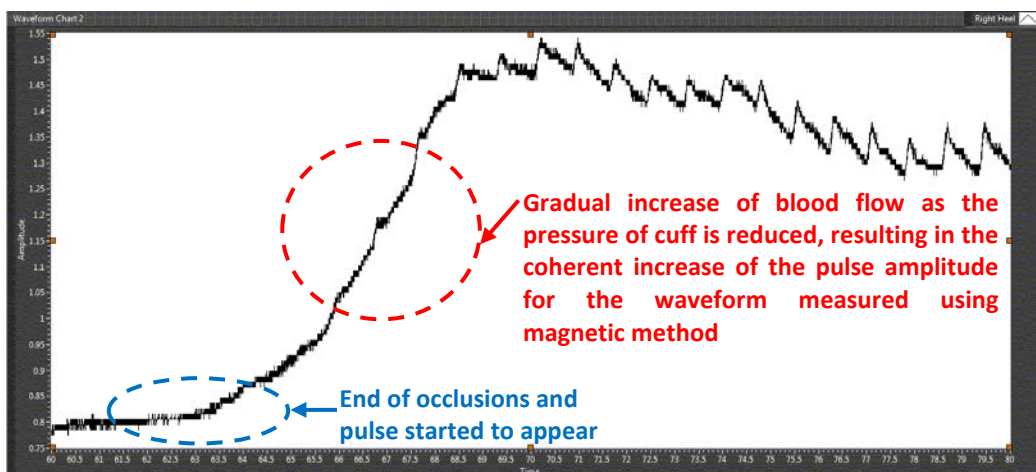


Figure 75 Waveform obtained using MMSB after occlusion

In order to compare the measured data quantitatively, the time to transit from occlusions to MF is measured (t_{meas}) for 6 healthy subjects, aged 20 to 24, and the results are tabulated in Table 15.

Based on the tabulated results, it can be observed that the measurements of PORH using MMSB are similar as those using the DRT4 instrument with less than 5% difference between the two measured values.

	DRT4 (Estimated)	MMSB (Actual)	% difference
Subject 1	5 sec	5.2 sec	4.00%
Subject 2	6 sec	6.1 sec	1.67%
Subject 3	5 sec	5.2 sec	4.00%
Subject 4	3 sec	3.1sec	3.33%
Subject 5	4 sec	4.1 sec	2.50%
Subject 6	5 sec	5.1 sec	2.00%

Table 15 Comparison of transit time from occlusions to MF for PORH measurements using MMSB and Moor’s Instrument, DRT4

6 Discussions of results

From the waveform obtained using MMSB for PORH, Figure 73, it can be observed that the occlusion period has no observable pulse waveform as blood flow is largely reduced resulting in non-observable magnetic disturbance. However, as the inflatable cuff releases its pressure at the end of the occlusion period, blood starts to flow with flow-rate relative to the rate of pressure released by the inflatable cuff. Therefore, the amplitude of the measured magnetic disturbance also increases with respect to the flow-rate. Therefore, from this experiment, it can be shown that MMSB is capable in measuring the changes in blood flow due to occlusion.

The experimental results obtained for PORH measurement using MMSB and laser Doppler are shown in Figure 71 and Figure 73 respectively. The similarity in the waveform characteristics in both figures for RF, occlusions and MF demonstrates the successful application of the MMSB method to acquire PORH data. This can be further compared using Figure 76 where both the MMSB and Laser Doppler data are normalized and plotted together. As shown, the characteristics response of a typical PORH measurement can be observed.

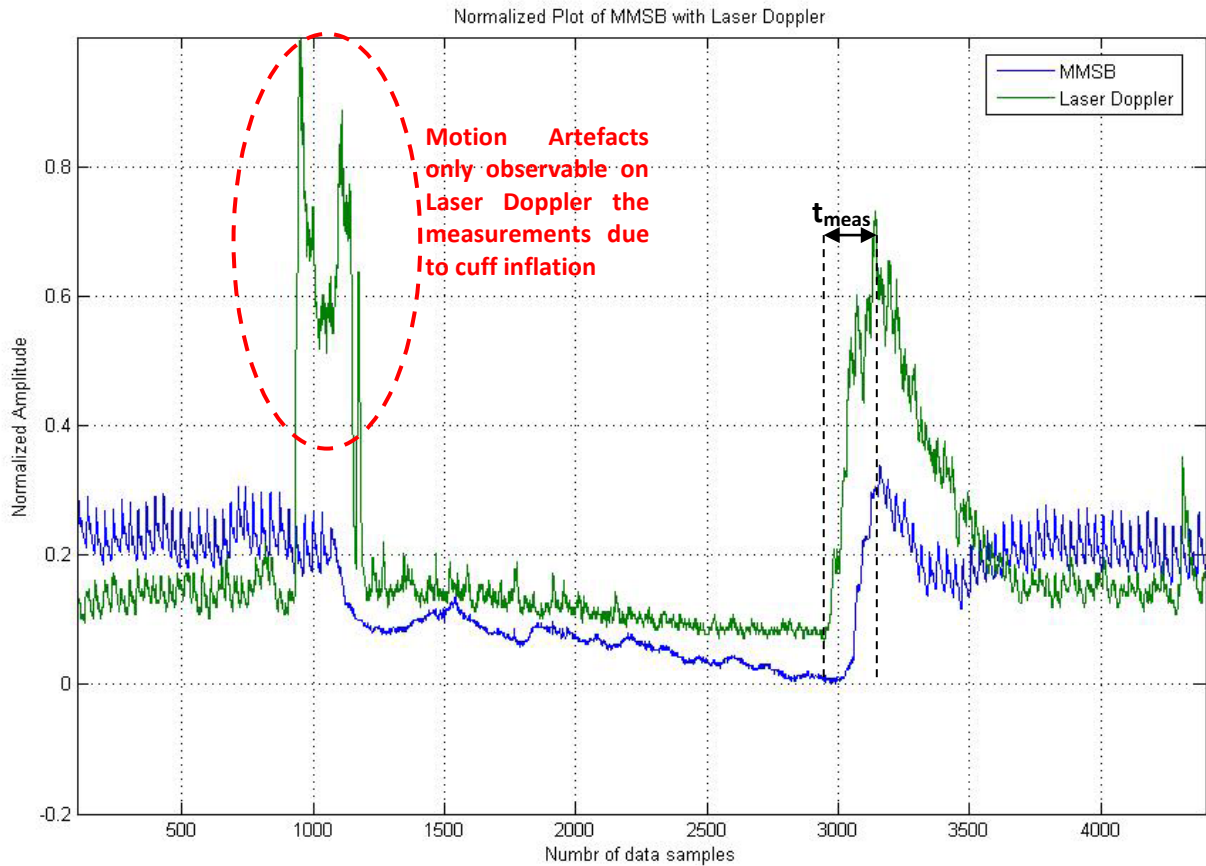


Figure 76 Typical normalized plot of MMSB with Laser Doppler measurements

In addition, the results obtained in Table 15, using the transit time from occlusion to MF as the reference, shows a difference of less than 5% in the t_{meas} between the MMSB method and the Laser Doppler method in acquiring the PORH data. Due to the data logging interval of 1 second provided by the laser Doppler equipment, the difference between these two methods of measurements can be accounted but not reducible. Therefore, the results obtained indicate the potential of using MMSB for PORH measurements.

7 Conclusions

The application of MMSB to acquire blood flow in the skin through the measurement on the change in amplitude of pulse signal at the end of the occlusion period is successfully demonstrated.

In addition, the PORH test has further supported the accuracy of MMSB through qualitative and quantitative analysis.

For qualitative analysis, the waveform obtained shows highly correlated results between laser Doppler and MMSB methods for signal acquisition of blood flow in the skin, where RF, occlusion and MF are clearly identified.

For quantitative analysis, the measured transit time from occlusions to MF for PORH measurements using MMSB and DRT4 for six healthy subjects have also proven the accuracy and reliability of the MMSB method for PORH measurements.

In conclusion, the application of MMSB for PORH measurements has proven to be a reliable and low cost approach in achieving PORH measurements. In addition, the benefit of non-invasive PORH measurement using MMSB method can provide useful alternative to clinical applications where good optical contact is not possible.

8 References

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Chapter 5: Measurement of blood pulse transit time using MMSB

1 Introduction

Use of cardiovascular pulsation, or commonly named as blood pulse, as one of the traditional indicators on the condition of the human body can diagnose different types cardiovascular diseases depending on the shape, amplitude and rhythm of this pulsation. Some of the possible diseases which can be diagnosed from different types of cardiovascular pulsation are listed in Table 16 [1]. A more detailed description of pulse shapes and the causes for abnormal pulse shapes are also listed in [2]. Factors that affect cardiovascular pulsation are the stroke volume of the heart, the compliance of the cardiovascular network and the character of the systolic heart ejection [1]. Different types of abnormal cardiovascular pulses are illustrated in Figure 77. From this, a sensor or sensing method that can detect the pulse, measure the pulse shape and the pulse transit time accurately; and further evaluate the elastic properties of the arterial wall, has great potential in medical and lifestyle applications.

Pulse Type	Physiological cause	Possible disease
Small and weak	Decreased stroke volume Increased peripheral resistance	Heart failure, hypovolemia, severe aortic stenosis
Large and bounding	Increased stroke volume Decreased peripheral resistance Decreased compliance	Fever, anaemia, hyperthyroidism, aortic regurgitation, bradycardia, heart block, atherosclerosis
Bisferiens	Increased internal pulse with double systolic peak	Aortic regurgitation, aortic stenosis and regurgitation, hypertropic cardiomyopathy
Pulsus alternans	Pulse amplitude varies from peak to peak, rhythms basically regular	Left ventricular failure

Table 16 Possible diseases which can be diagnosed based on the different types of cardiovascular pulse shapes [1]

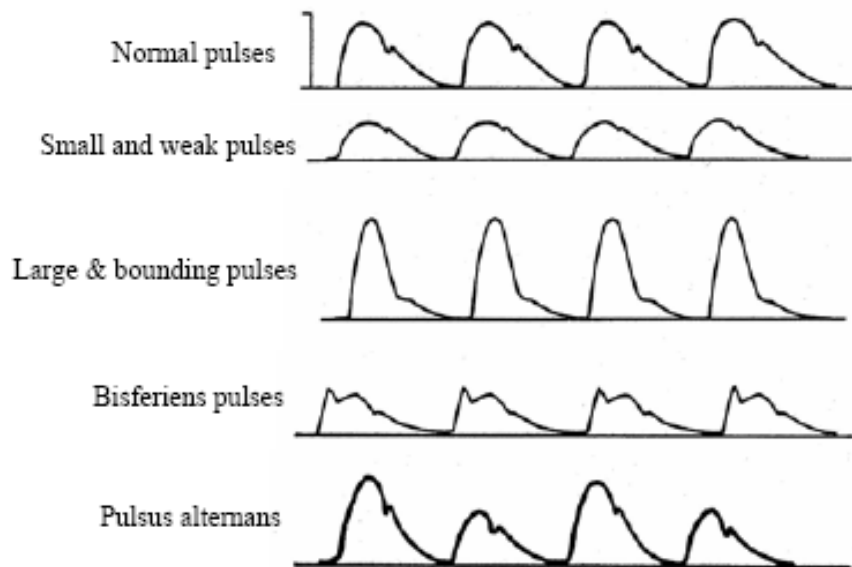


Figure 77 Different types of cardiovascular pulse shapes [1]

To study the cardiovascular pulsation, blood pressure is one of the widely used indicators in both medical and lifestyle applications. With the increase in popularity of blood pressure measurement devices in the last decade, prices of these measurement devices have sunk to an appropriate level for ordinary consumers. The incorporation of automatic measurement features and ease of use have also contributed to the growing popularity of blood pressure measurement devices as a lifestyle devices. However, such measurement devices are typically cuff based and rely on indirect Blood Pressure (BP) measuring method using the detection of Korotkov-sounds [3]. This method had been used for more than 100 years and has two major deficiencies. First, the cuff grossly intervene the measured parameter and second, it determines the pressure only at a single point of time.

Oscillometric method [4], applied in automatic home BP-meters, apart from the aforementioned insufficiencies has also a great built-in uncertainty: it empirically calculates the systolic and diastolic value from the measured mean pressure.

Tonometry [5] is, however, a very different approach. Although a pressure is applied to the artery to press it against the bone, the pressure is not as strong as to completely occlude the artery. Rather, the pressure is maintained at a level that counteracts the circumferential tension of the arterial wall. Therefore, it is sometimes named the “vascular unloading” technique [6]. Since tonometry is to be applied to the peripheral circulation, an algorithm is often used to estimate arterial BP from the recorded pressure.

Although these automatic BP (ABP) meters can provide objective BP readings, all of them, like the conventional mercury sphygmomanometer, incorporate the use of a cuff in their working principles.

In addition, the size of the cuff must not be too small since overestimation of BP by using an inappropriately small cuff has been well documented [7]. Thus, this renders the reduction of size, cost and power consumption in these devices very difficult. Moreover, the occlusion of the artery increases the workload of the heart and causes circulatory interference at the measurement site. Therefore, they are not preferred to be used frequently. Compared to the cuff-based approaches, a cuff-less technique using pulse transit time is more desirable [8].

Researchers have long reported that pulse transit time (PTT) is directly related to ABP and can be used as an index for the beat-to-beat BP estimation [8]-[11]. Typical PTT is the measure using the time difference between the R-wave peak of electrocardiogram (ECG) and maximum slope of the finger photoplethysmogram (PPG) as shown in Figure 78. However, the use of the finger PPG will require continuous optical contact with the skin on the finger and is restrictive for daily activities as the finger is typically covered for optimal signal acquisition by the PPG [12].

In this chapter, the magnetic method of blood pulse acquisition is applied on the wrist in place of the finger PPG leading to the development of a device that is capable of continuous PTT measurement leading to the measurement of ABP in an unrestrictive manner suitable for daily activities.

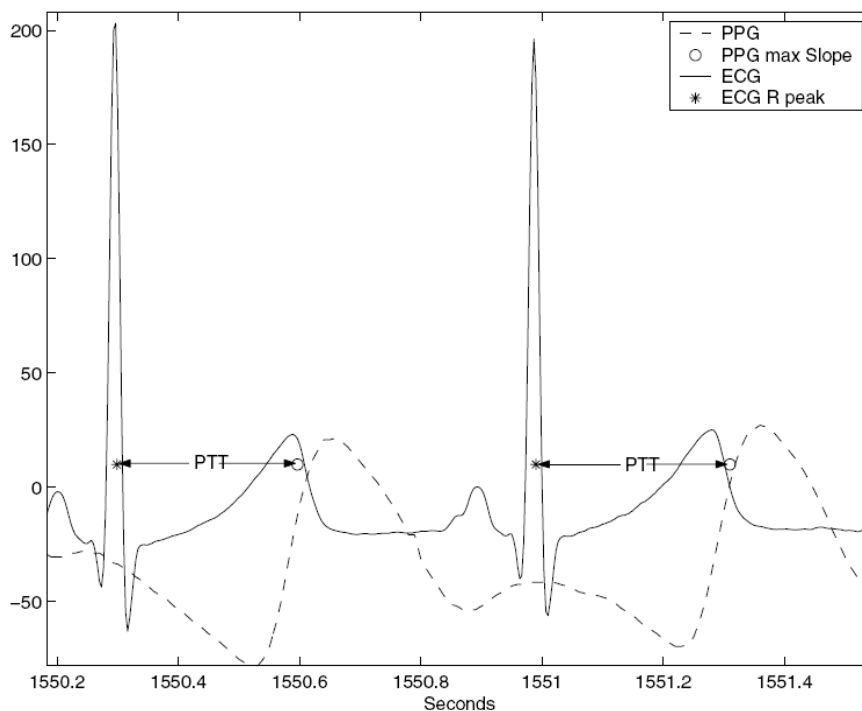


Figure 78 Illustration of PTT Definition [6]

2 Setup for blood pressure measurement using oscillometric based BP meters

There are many commercially available cuff based blood pressure monitoring devices which are designed for lifestyle applications. In this experimental setup, the OMRON Intellisense Blood Pressure Monitor - Arm Model IA1/IA1B is used for acquisition of blood pressure using the setup as shown in Figure 79.

Using this setup, 10 young subjects aged 21 to 25 are recruited as voluntarily test subjects for this experiment. The results obtained from this measurement are shown in Table 17. The measured Systolic (SYS) and Diastolic (DIA) pressure can be associated to the heart activities where during heart contraction, the blood pressure in the artery increases and diastolic blood pressure changes to systolic blood pressure. The difference of these pressures is called pulse pressure. Normal blood pressure occurs when systolic pressure is less than 130 mmHg and diastolic blood pressure is less than 80 mmHg for adults [1].

Mean arterial pressure (MAP) is the mean pressure over one cardiac cycle. Using the Equation 61 [13], the MAP can be calculated and is tabulated in Table 17 for comparison with measurements in the next section.

Equation 61
$$MAP = P_{DIA} + \left(\frac{P_{SYS} - P_{DIA}}{3} \right)$$

where P_{DIA} = Diastolic pressure in mmHg

P_{SYS} = Systolic pressure in mmHg

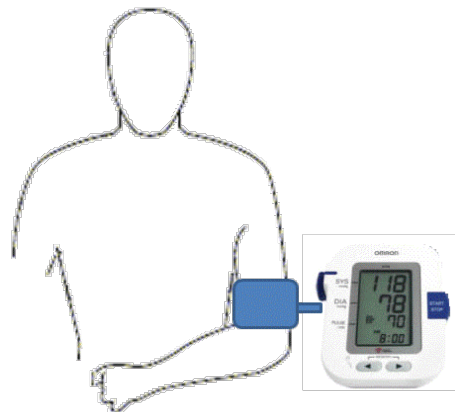


Figure 79 Experiment setup using OMRON Intellisense Blood Pressure Monitor - Arm Model IA1/IA1B

<i>Name</i>	<i>Distance from heart to wrist (cm)</i>	<i>Pulse Transit Time (sec)</i>	<i>MAP (mmHg)</i>
Subject 1	81.0	0.22	86.67
Subject 2	75.0	0.21	81.33
Subject 3	77.0	0.21	84.67
Subject 4	85.0	0.22	91.33
Subject 5	75.0	0.22	75.67
Subject 6	74.0	0.20	86.33
Subject 7	78.0	0.21	86.67
Subject 8	71.0	0.17	113.67
Subject 9	84.0	0.21	97.00
Subject 10	77.5	0.22	76.67

Table 17 Measurement results for BP and MAP using BP-meter

3 Experimental setup for blood pressure measurement using MMSB

The experimental setup is shown in Figure 80 where the signals acquired from the ECG electrodes and MMSB are concurrently being sampled and processed using the gTEC USBamp. The data post-processing is completed in MATLAB Simulink and the typical waveform obtained is shown in Figure 81.

Using Figure 81, the PTT is measured from the peak of the R-wave to the maximum slope of the MMSB signal and tabulated in Table 18. In addition, the distance between the heart and the wrist (where the MMSB signal is acquired) is also measured as illustrated in Figure 82 and tabulated in Table 18. The pulse wave velocity (PWV) of blood is calculated based on the measured distance and PTT values. From research data available [2], typical values of PWV range from 5 m/s, with young people in central arteries, to 15 m/s, in peripheral and old arteries. Using this as the basis of comparison, the cardiovascular performance of the 10 young subjects is found to be within the acceptable range of 5m/s confirming the measured values for PTT and distance between heart and wrist is reasonable.

The work done by the pulse wave can be expressed in terms of the kinetic energy of the wave and the gravitational potential energy as shown in Equation 62.

Equation 62
$$F \cdot d = \frac{1}{2}mv^2 + mgh$$

where F = force exerted on blood
d = distance from heart to wrist (or fingertip)
m = mass of blood
v = pulse wave velocity
g = 9.81m/s²
h = height difference between heart and wrist (or fingertip)

The force exerted on blood can also be written in terms of pressure difference as shown in Equation 63:

Equation 63
$$F = \Delta BP \cdot a$$

where a = cross section area of the artery
 ΔBP = pressure difference between the two sites, the heart and the wrist or fingertip

Substituting Equation 62 into Equation 63 and rearranging the variables:

Equation 64
$$\Delta BP = \frac{1}{2} \frac{m}{a \cdot d} v^2 + \frac{m}{a \cdot d} gh$$

Replacing $\frac{m}{a \cdot d} = \rho$ (density of blood) and $v = \frac{d}{PTT}$ (pulse wave velocity of blood), Equation 64 can be expressed as shown in Equation 65:

Equation 65
$$\Delta BP = \frac{1}{2} \rho \frac{d^2}{PTT^2} + \rho gh$$

The distance d can be approximated using measurements of the distance between subject's heart and wrist as illustrated in Figure 82. In addition, PTT is the measured pulse transit time in seconds with reference to the ECG signal as illustrated in Figure 78. Furthermore, the average blood density ρ is reported to be 1050 kg/m³ [14]. Therefore, using the values obtained for these parameters, ΔBP in Equation 65 can be numerically estimated.

Based on research findings [15], it had been reported that the pressure drop in the arterial side of circulation accounts for roughly 70% of the total pressure drop in the body. Therefore the subject's average change in the blood pressure or mean arterial pressure, MAP, can be approximated using Equation 66[15].

Equation 66
$$MAP = \frac{1}{0.7} \left(\frac{1}{2} \rho \frac{d^2}{PTT^2} + \rho gh \right)$$

- where MAP = mean arterial pressure
 ρ = density of blood, 1050 kg/m³
 g = 9.81m/s²
 h = height difference between heart and wrist (or fingertip)
 d = distance from heart to wrist (or fingertip)
 PTT = Pulse transit time

Using the Equation 66 derived from the basic principle, the MAP can now be calculated using the measured value of PTT from the MMSB method of pulse acquisition.

For the relative height parameter, h in Figure 80 and appears in Equation 66, the values ranges between 7cm to 9cm for all the subjects in this experiment. As such, the mean value h = 8cm is adopted for all ten subject.

In addition, the adaptations of the mean value for h aims to test the robustness of the measurement method to facilitate future implementation of the MMSB method, for blood pressure measurements, as a lifestyle device. However, the use of a mean value of h is found to induce additional error, which will be explained later in this chapter (Reference Table 19)

The calculated MAP values are tabulated as shown in Table 18. The results of this calculation for MAP will be compared using the values obtained from the use of the commercial BP-meter using Equation 61. The accuracy of the use of the PTT obtained using MMSB to measure MAP is analyzed and compared with the commercial BP-meter. The results will be presented and discussed in the next section.

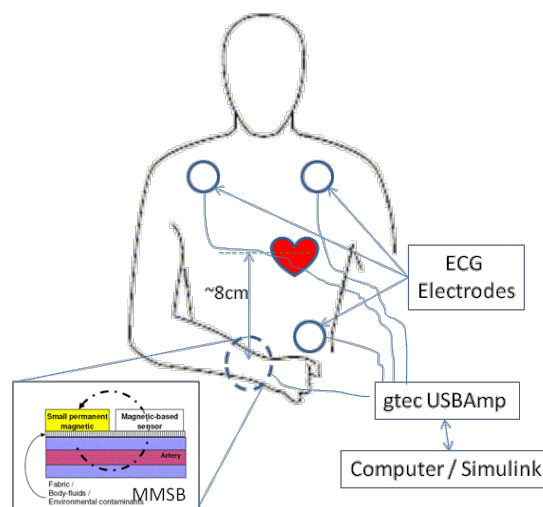


Figure 80 Experimental setup used for PTT measurements using MMSB and ECG

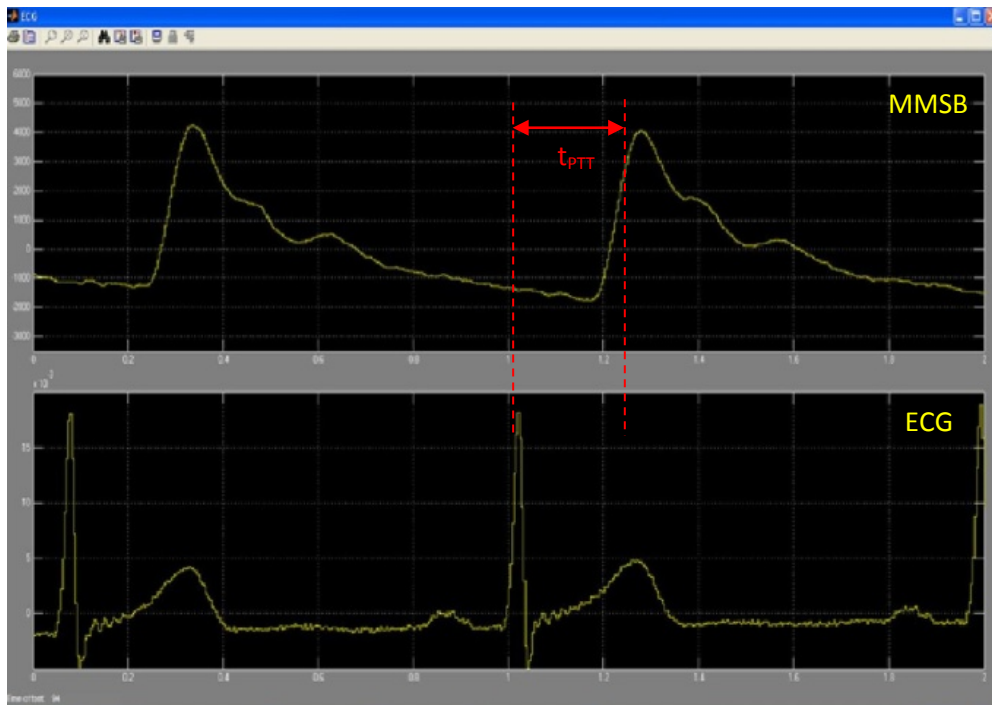


Figure 81 Waveform for MMSB and ECG signal with PTT measurement

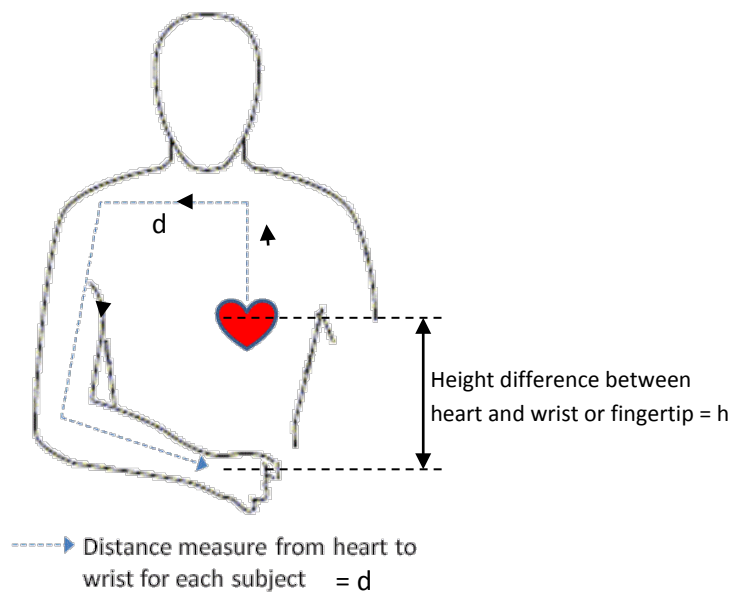


Figure 82 Illustrations of distance measurements from heart to wrist

<i>Name</i>	<i>Distance from heart to wrist (cm)</i>	<i>Pulse Transit Time (sec)</i>	<i>PWV (m/s)</i>	<i>Estimated MAP (mmHg)</i>
Subject 1	81.0	0.220	3.68	84.00
Subject 2	75.0	0.210	3.57	79.49
Subject 3	77.0	0.215	3.58	79.89
Subject 4	85.0	0.225	3.78	88.03
Subject 5	75.0	0.225	3.33	70.24
Subject 6	74.0	0.190	3.89	93.07
Subject 7	78.0	0.200	3.90	93.30
Subject 8	71.0	0.170	4.18	105.87
Subject 9	84.0	0.210	4.00	97.44
Subject 10	77.5	0.230	3.37	71.61

Table 18 Measurement results for distance from heart to wrist, PTT and PWV with MAP estimated using MMSB measured data

4 Discussions of results

Using the measured and estimated MAP from Table 17 and Table 18 respectively, the results obtained for this experiment can be compared as tabulated in Table 19 and plotted as shown in Figure 83.

By the calculating the percentage difference between measured and estimated MAP, it can be observed from Table 19 that the difference in both MAP data is less than 8%. This difference can be accounted for in the MAP estimation due to the introduction of uncertainty by the values for d , h and PTT in the equation through estimation measurements.

In addition, h was assigned as a constant in the estimation of blood pressure which may have potentially increased the uncertainty of MAP. This could have resulted in the difference of MAP values obtained using the two methods.

In order to estimate the uncertainty contributed by each of variables in Equation 66, the uncertainty for each variable can be determined from Equation 66 by assuming that ρ , h and g are constants (i.e. not used in uncertainty propagation). Using the uncertainty formula in Equation 67 for product of factors raised to powers, Equation 66 can be simplified as Equation 68.

Equation 67 $f = x^m y^n \rightarrow \varepsilon_f = \sqrt{m^2 \varepsilon_x^2 + n^2 \varepsilon_y^2}$

Equation 68 $\varepsilon_{MAP} = \sqrt{2^2 \varepsilon_d^2 + (-2)^2 \varepsilon_{PTT}^2} = 2 \sqrt{\varepsilon_d^2 + \varepsilon_{PTT}^2}$

The uncertainty contributed by the distance (i.e. ε_d) and PTT measurement (i.e. ε_{PTT}) is estimated as 2% each due to the measurement inaccuracy. This will result in the total amount of uncertainty to be estimated as 5.66%. Taking into the account the parameter h is not a constant; the difference between the estimated uncertainty of 5.66% and the maximum measured error of 8% in Table 19 can be accounted by taking into consideration the uncertainty contributed by the measurement parameter h.

The objective of proving the feasibility of using MMSB for PTT measurements for MAP measurements has been successfully demonstrated. The results obtained in this experiment are encouraging as the measurement setup has proven the feasibility potential for the applications of MMSB in place of PPG for PTT measurements. Most important of all, the use of MMSB will allow the PTT signal be acquired on the wrist resulting in minimum inconvenience for execution of daily activities.

<i>Subjects</i>	<i>Measured MAP (mmHg)</i>	<i>Estimated MAP using PTT measured from MMSB method (mmHg)</i>	<i>% Difference between measured and estimated MAP</i>
Subject 1	86.67	84.00	-3.08%
Subject 2	81.33	79.49	-2.26%
Subject 3	84.67	79.89	-5.65%
Subject 4	91.33	88.03	-3.61%
Subject 5	75.67	70.24	-7.18%
Subject 6	86.33	93.07	7.81%
Subject 7	86.67	93.30	7.65%
Subject 8	113.67	105.87	-6.86%
Subject 9	97.00	97.44	0.45%
Subject 10	76.67	71.61	-6.60%

Table 19 Tabulated results for measured MAP using BP meter with estimated MAP using PTT with MMSB measurements

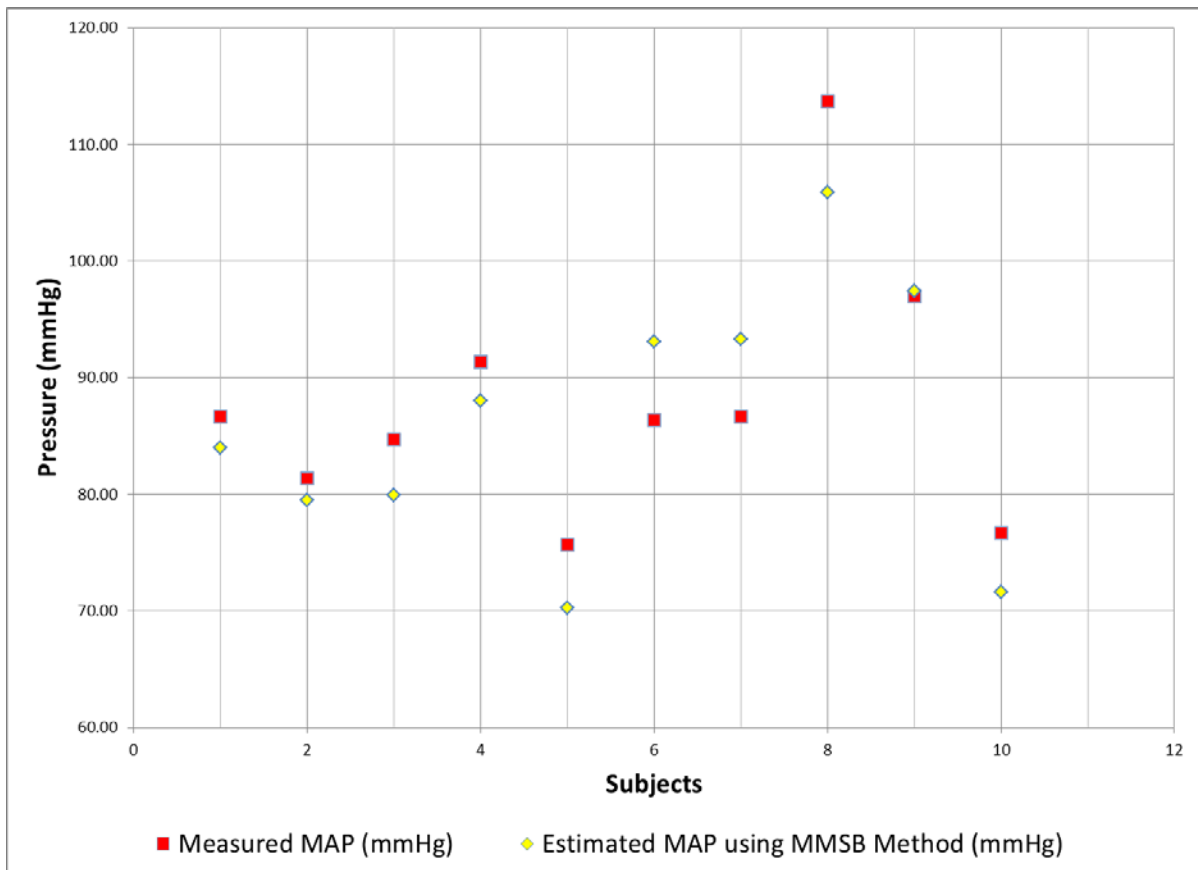


Figure 83 Plot of Measured MAP and Estimated MAP using PTT measured from MMSB method

5 Conclusions

The application of MMSB to acquire PTT at the wrist in place of the PPG at the finger is more convenient for the continuous monitoring of physiological signs such as the blood pressure. The result of such an acquisition is qualitatively and quantitatively assessed using the waveform (Figure 81) and MAP measurement (Table 19) respectively. From the waveform, the ease of identifying the measurable PTT value with respect to PPG method is illustrated. In addition, the measured and calculated MAP data shows good correlations with errors of less than 8% with respect to the measurements obtained from the use of the lifestyle OMRON BP-meter. Therefore, it can be concluded that the use of MMSB for PTT measurements leading to the determination of the MAP has been successfully demonstrated.

With the completion of MMSB for blood pulse transit time indicating the measurable parameter, MAP, the following chapter will focus on the development of devices based on MMSB for heart rate and relaxation analysis.

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Chapter 6: Applications of MMSB

1 Introduction

The design and development of the MMSB in a laboratory setup has been proven in the previous chapters. In this chapter, two applications of the MMSB will be developed to achieve an advanced next generation bio-monitoring of physiological vital signs. To achieve this objective, a wireless deployable platform for pulse rate sensing will be developed using the wireless ad-hoc network for concurrent and continuous monitoring of blood pulse in a multi-user environment. This development is termed “Wireless Pulse Sensing” and in this development, a wearable low power wireless system will be developed with signal acquisition for remote monitoring of pulse rate. This is followed by the development of a standalone pulse rate monitoring device with LCD for local display and self-monitoring.

In the second development for this chapter, MMSB will be integrated into an existing lifestyle product for relaxation assessment using blood flow information collected before and after massage. In this section, the project will be termed “Relaxation assessment using MMSB” where algorithms were developed to statistically compare the changes in blood flow before and after a massage. The changes in blood flow and heart rate variability (HRV) will be used to assess the mental state of being stressed or relaxed for the subject based on published results.

Finally, this chapter will conclude on the results obtained from these two developments and discuss on future development of applications of MMSB.

2 Wireless Pulse Sensing

As discussed in Chapter 1, there are many commercially available heart rate measurement devices and systems where each of these solutions has its associated advantages and disadvantages. The advantages of using MMSB as the pulse sensing method to measure heart rate has been comprehensively presented in this dissertation. This section will describe the integration of MMSB with a wireless ad-hoc network as a wearable device on the wrist to continuously monitor heart rate of subjects. This work is funded by the Totaliser Board of Singapore at S\$350,000.

2.1 Introduction

The wireless ad-hoc network required to support this development work has to be low power and scalable for applications such as mass run and military trainings. Therefore, the various wireless ad-hoc networks available are reviewed and the 2.4GHz ZigBee / IEEE 802.15.4 standard was selected due to its low power and scalability.

With the network protocol selected, existing commercial products that provide the ZigBee solution were reviewed and the XBee and XBee-PRO ZigBee™ 802.15.4 OEM RF Modules by MaxStream, Inc. is found to be the most suitable due to the followings:

1. Low cost < USD\$21
2. Small footprint – 2.438 cm x 2.761 cm
3. Scalability – able to support multiple users with ad-hoc network forming and hopping
4. Ease of integration – integrated 10 bit ADC; on-board chip antenna; data interface available
5. Wide operating input voltage range – 2.1-3.6V
6. Low current – transmit current of 35mA; receive current of 40mA; sleep current <1 μ A at 25°C
7. RF range – Indoor/Urban up to 133 ft (40 m); Outdoor/RF line-of-sight up to 400 ft (120 m)
8. Configurability of the RF modules – same RF module can be configured as repeater or base or remote device with 65,000 network addresses available for each channel

The intended use of the ZigBee™ based wireless ad-hoc network is to support the concurrent monitoring of multiple users, where the wireless network is to be configured for Point-to-Point, Point-to-Multipoint, Peer-to-Peer and Mesh configurations. Based on this intent, the system overview of the final design system is shown in Figure 84 where base station can remotely receive data (pulse rate) through the ad-hoc mesh network automatically formed during deployment.

For the purpose of this development, the data acquired by the ADC is wirelessly transmitted back to the base station for processing by the PC using the graphical development tool, LabView 8.0 from National Instruments. Upon successful implementation of the PC based processing, the algorithm developed will be implemented onto an embedded processor with LCD display to function as a standalone wearable pulse rate monitoring device.

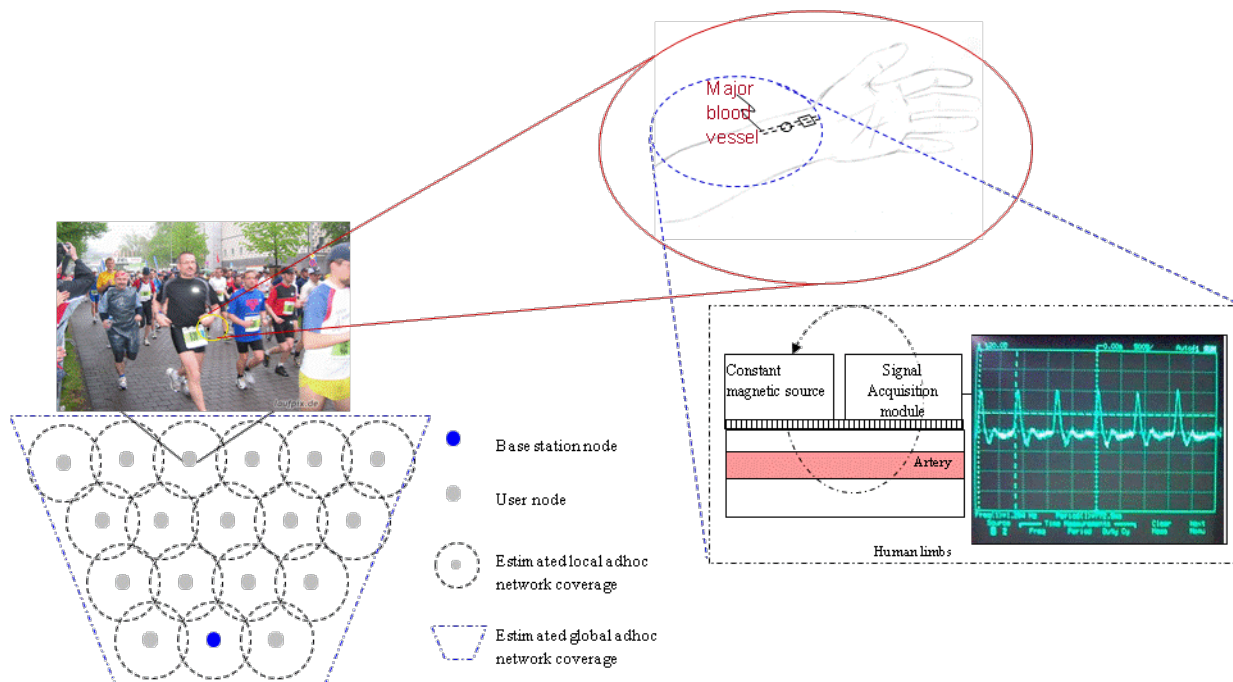


Figure 84 Illustration of wireless pulse sensing system for monitoring of heart rate using MMSB

2.2 System design

The system architectural design can be classified into two main frameworks as illustrated in Figure 85. The framework on the right of Figure 85 will address the many challenging aspects of the development of user node with wireless interface to support multi-user environment where a significant amount of work is devoted to the configuration of the ZigBee™ 802.15.4 OEM RF Modules by MaxStream, Inc. to support the wireless ad-hoc network and the software development for the base station node to support concurrent monitoring of multiple users. The framework on the left of Figure 85 will focus on the end-user node development which includes the MMSB signal conditioning, data acquisition and network interface.

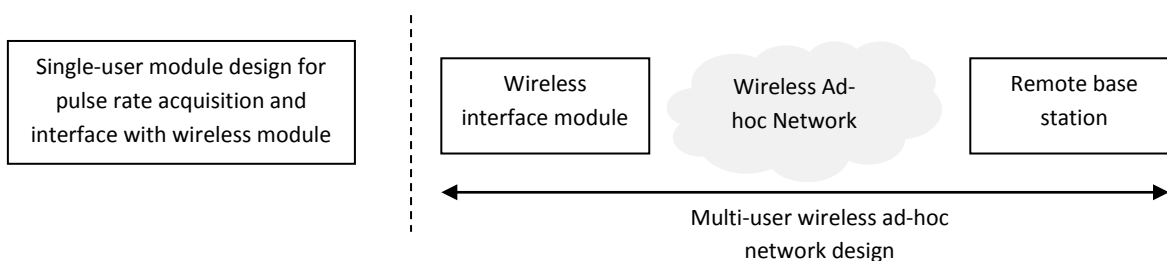


Figure 85 Illustration of System Architecture of Wireless Pulse Sensing System

2.2.1 Signal Conditioning

The MMSB signal derived from the GMR based magnetic sensor is typically 0.2 mV peak-peak. Therefore, an amplification of about 100 times is necessary to render this signal usable for signal processing via a 10 bit Analog-to-Digital Converter (ADC) to achieve heart-rate detection. Realizing clean amplification of the MMSB signal with such high gain is no easy task given that the human body moves and motion artifacts causes a lot of noise to the MMSB sensing design. As such, the MMSB signal has to be filtered by a strong post filter after amplification. Unfortunately, any amplification will also amplify the noise voltages in addition to the desired MMSB signal. In certain situations, the noise can completely override the MMSB signal and render the amplified signal useless.

In order to minimize the effects of common mode noise due to motion artifacts, the use of differential signal conditioning is implemented. Since the output of the GMR sensor is based on the differential output of a Wheatstone bridge, an ideally matched differential amplifier is selected where common mode noise from the GMR sensor can be automatically cancelled out. Therefore, the differential amplifier used in the front end of this application is the INA118 instrumentation amplifier (Reference ANNEX A) that has perfectly matched and balanced integrated gain resistors.

The final signal conditioning circuit used for this development is shown in Figure 86, where the output voltage can be described using Equation 69. The high-performance Instrumentation Amplifier (IA) INA118 has greatly simplified the design of bridge preamplifiers while adding significant advantages in noise, size, and performance over op amp implementations.

$$\text{Equation 69} \quad V_{out} = \left(1 + \frac{50K}{R_G}\right) (V_{out+} - V_{out-})$$

where the frequency 3dB point is given by $f = \frac{1}{2\pi R_3 C_3}$

The frequency response of circuit is designed to acquire the heart rate (typically around 1Hz). However, due to the low amplitude of the MMSB signal, the filter was found to be inappropriate as it will attenuate the desired signal significantly. As such, the components R_3 and C_3 are bypassed, where the MMSB signal is directly amplified and sampled by the ADC for wireless transmission.

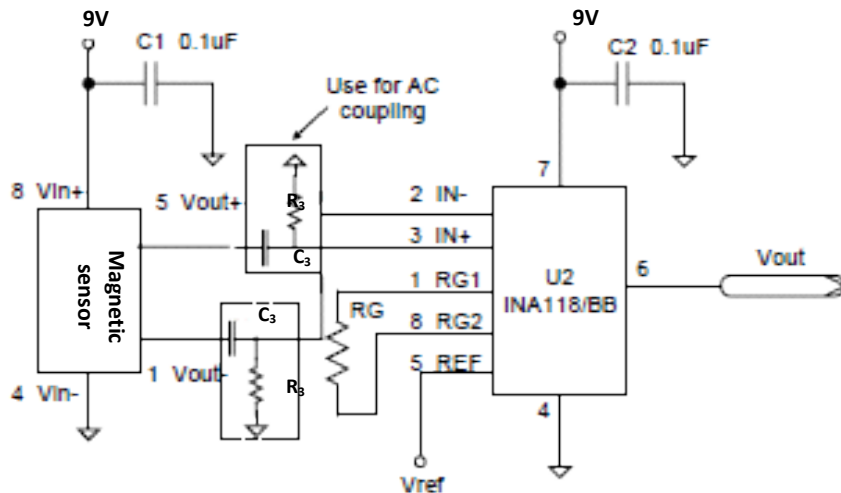


Figure 86 Signal conditioning circuit WiPS

2.2.2 Analog-to-Digital converter

The XBee RF Modules used for this development has an integrated support for a 10 bit Analog-to-digital conversion (ADC) and is sharing similar ports for the digital Input / Output (I/O) line passing as shown in Table 20. The XBee has 6 usable A/D pins that will allow the user to utilize one (sampling rate of up to 1KHz) or all (assuming sufficiently slow sample rates when all ADCs are used i.e. 1KHz/6 = 167Hz) of these ADCs. However, in this development, only one ADC channel will be used at a sampling rate of 10Hz to minimize bandwidth required for transmission of ADC data. The signal processing for heart rate measurements will be done at the base-station so as to simplify the design of the wearable device and reduce the power consumption.

AD = Analog-to-Digital Converter, DIO = Digital Input/Output
 Pin functions not applicable to this section are denoted within (parenthesis).

Pin Function	Pin#	AT Command
AD0 / DIO0	20	D0
AD1 / DIO1	19	D1
AD2 / DIO2	18	D2
AD3 / DIO3 / (COORD_SEL)	17	D3
AD4 / DIO4	11	D4
AD5 / DIO5 / (ASSOCIATE)	15	D5
DIO6 / (RTS)	16	D6
DIO7 / (CTS)	12	D7
DIO8 / (DTR) / (Sleep_RQ)	9	D8

Table 20 Pin functions and associated pin numbers and commands for the XBee RF Modules

Using Table 20, the ADC channel is configured to sample the analog signal at 10 Hz and store the sampled signal as a digital format for transmission over the wireless network. With the XBee

integrated ADC, it will simplify the circuit design allowing the XBEE wireless module to be connected directly to the output of the amplifier where data will be sent in a serial form, wirelessly to the base station.

In order to minimize power due to RF transmission, the buffering feature of the XBee module is also implemented. The XBee module has a total buffer of up to 93 bytes and using this buffer size, it will allow the user to set the number of samples stored before the data is transmitted. With the 10 bits ADC integrated in the XBee module, each sample will utilize two bytes. This will lead to a maximum buffer size of 46 samples before transmission. With one ADC configured to operate at a sampling rate of 10Hz (i.e. 10 times over sampling for typical heart rate of 1Hz), data is configured to be buffered up to 4 seconds before each RF transmission. This will allow significant power saving as compared to continuous monitoring as shown in the reduction of average current in Table 21.

Scenarios		A	B	C
Time	Units	Scenario_A	Scenario_B	Scenario_C
Sleep	s	0	2	4
Idle/Receive	ms	40	40	40
Transmit	ms	3.332	4.74	6.212
Radio type		XBee	XBee	XBee
# of bytes transmitted		2	46	92
Total System Current				
Sleep	mA	0.002	0.002	0.002
Idle/Receive	mA	45	45	45
Transmit	mA	40	40	40
Power usage comparison				
Sleep	%	0.00%	0.20%	0.39%
Idle/Receive	%	93.11%	90.29%	87.53%
Transmit	%	6.89%	9.51%	12.08%
Average current	mA	44.6155266	0.9749895	0.5082482

Table 21 Estimation of average current for Scenarios A (unbuffered), B (2 seconds buffer) and C (4 seconds buffer) using the power estimator provided for XBEE transceiver

2.2.3 Data interface to wireless module

Once the data buffer is full, the XBEE will package the ADC data with the data overhead, such as the delimiter, length of bytes and source address, for transmission over the RF channel. This is automatically done by the integrated processor within the XBEE module and the base-station will have to decode the ADC data for signal processing to calculate heart rate. A typical data packet sent over the RF channel with the buffer size set to 2 is illustrated below.

7E 00 10 83 56 78 22 00 05 06 00 **00 00 03 FF** 77

where the serial data stream can be broken down as:

7E Start Delimiter
00 10 Length Bytes
83 API Identifier Byte for 16bit A/D data (82 is for 64bit A/D data)
56 78 Source Address Bytes
22 RSSI Value Bytes
00 Option Byte
05 Sample Quantity Byte
06 00 00000110 00000000 Channel Indicator *
00 00 1st Sample Data ADC (min value for ADC is 00 00)
03 FF 2nd Sample Data ADC (max value for A/D is 03 FF)
77 Check Sum

2.3 Data processing for heart rate measurements

The sampled MMSB waveform is transmitted over the RF channel to the base-station PC for signal processing. Data is received using the ZigBee receiver and processed using the LabView programming interface where the pulse rates for multiple users are measured and displayed as shown in Figure 87 and Figure 88 respectively.

In Figure 87, data received from the ZigBee module is first processed to extract the header and concatenate the data together. The concatenated data will be passed to the pulse rate measurements module as shown in Figure 87.

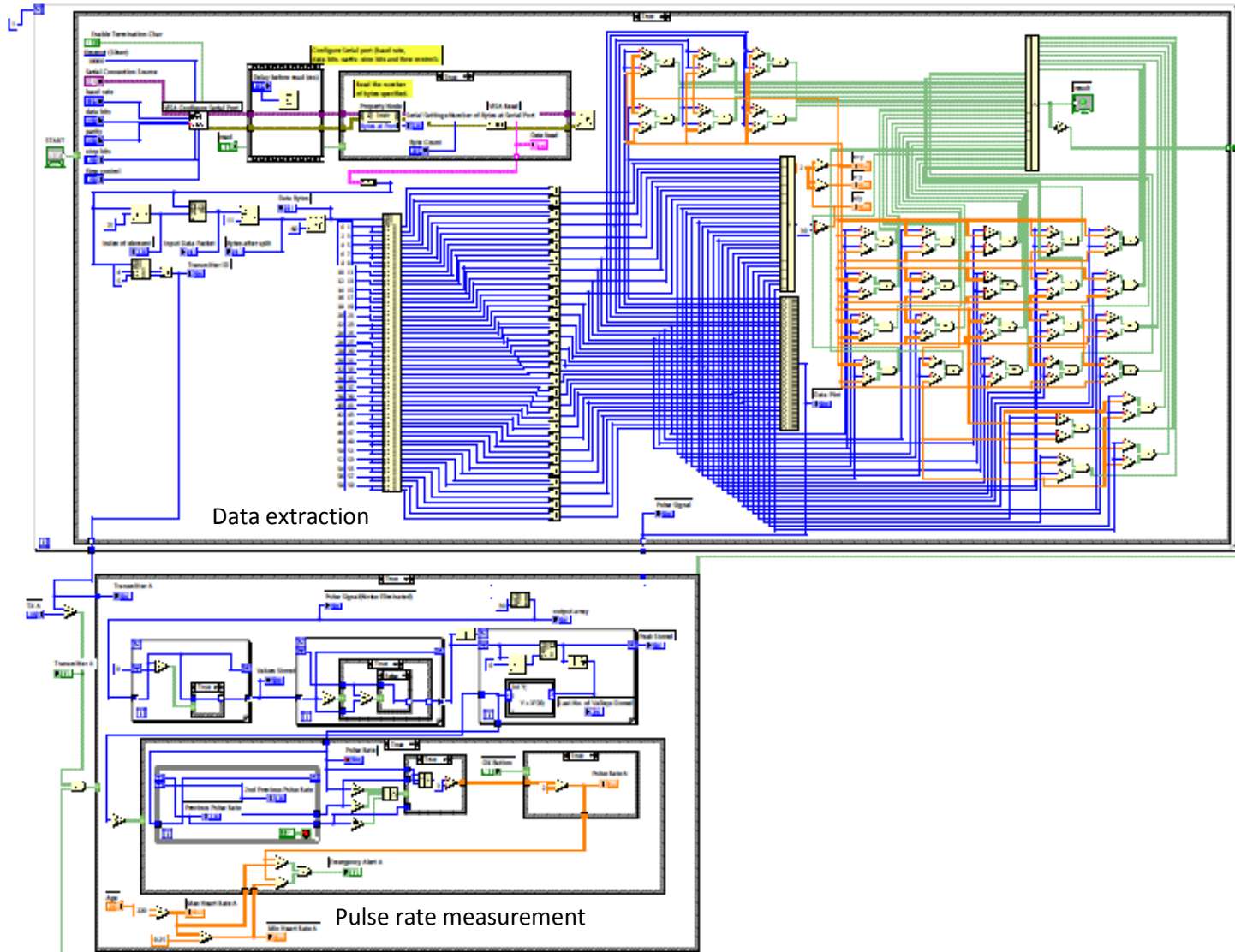


Figure 87 Program developed using LabVIEW to extract data from the ZigBee receiver and processed for pulse rate measurements

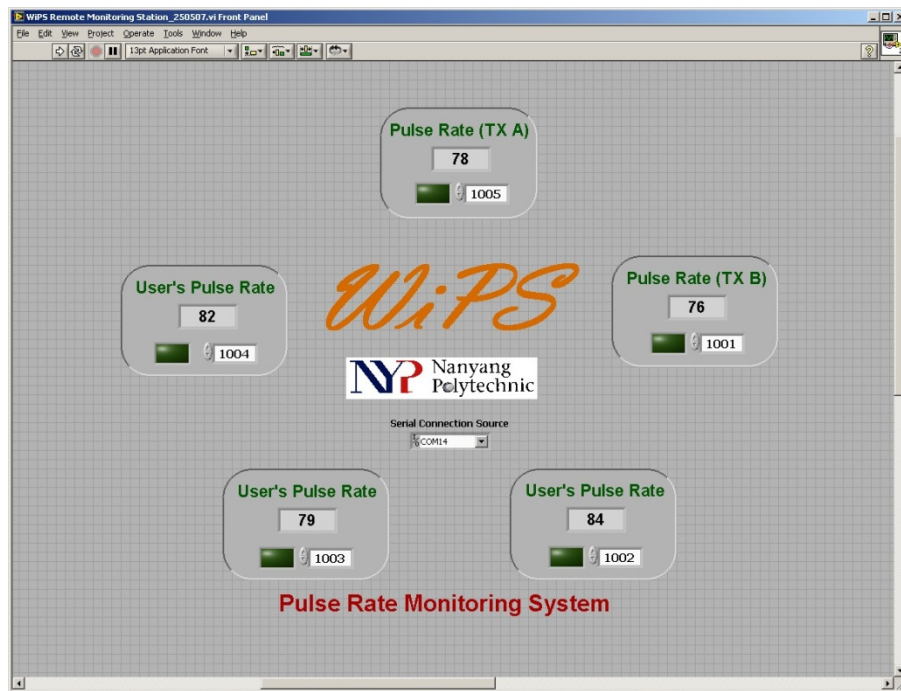


Figure 88 Concurrent display of pulse rate for multiple users

In order to measure the pulse rate, it is important to detect the peak of the MMSB signal. As such, a two stage filtering architecture is used in Matlab / Simulink model as shown in Figure 89. Based on Figure 89, the high frequency noise in the MMSB signal is first removed through the implementation of a digital low-pass FIR filter. This is followed by differentiating the low-pass filtered samples of the MMSB signal using a high-pass filtered so as to isolate the peak of the MMSB waveform. Finally, the output of the peak detected MMSB waveform is passed through a zero crossing module for heart rate measurements. The model is verified through the waveforms obtained in Figure 95 where the peak is accurately detected for each heart beat.

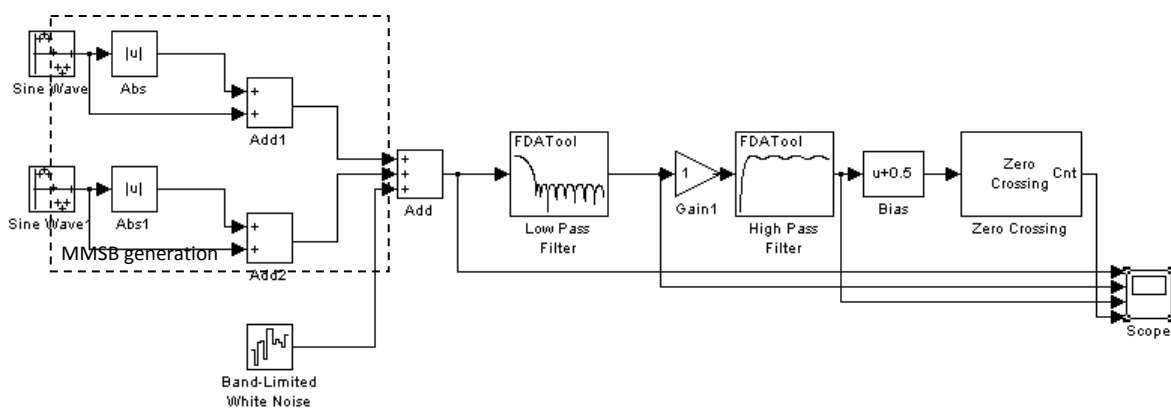


Figure 89 Architecture of filter design peak discrimination

The low-pass filter is designed based on a N-tap FIR filter using the difference equation of the form shown in Equation 71 [1].

Equation 70 $y(n) = \sum_0^{N-1} h(k)x(n - k)$

where $h(k)$ is the impulse response of the filter

Equation 71 maps directly into the direct form FIR structure as shown in Figure 90. Using the FIR structure and that the typical frequency of human heart activity being less than 3Hz, the filter was designed to have a pass-band upper frequency of 6 Hz and stop-band lower frequency of 30 Hz. Using the filter design toolbox from the mathematical software MATLAB, the frequency response of the filter is designed as shown in Figure 91 where a 17-tap FIR direct form filter with equiripple was generated. In order to provide additional gain for the MMSB signal, the filter coefficients are scaled to compensate for the filter attenuation and the magnitude response of the filter is illustrated in Figure 92. Therefore, at the digital filter output of the signal processor, the total amplification factor provided for the MMSB signal with respect to the noise can be greater than 200 times (i.e > 46 dB of voltage gain from filtering).

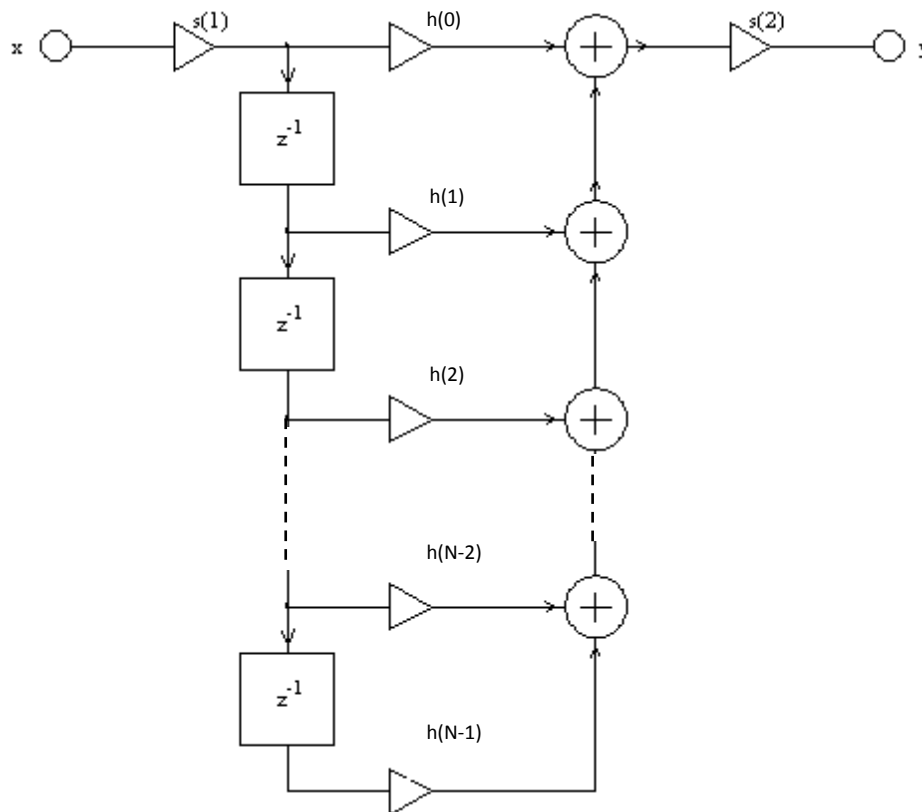


Figure 90 Illustration of a typical Direct Form FIR filter

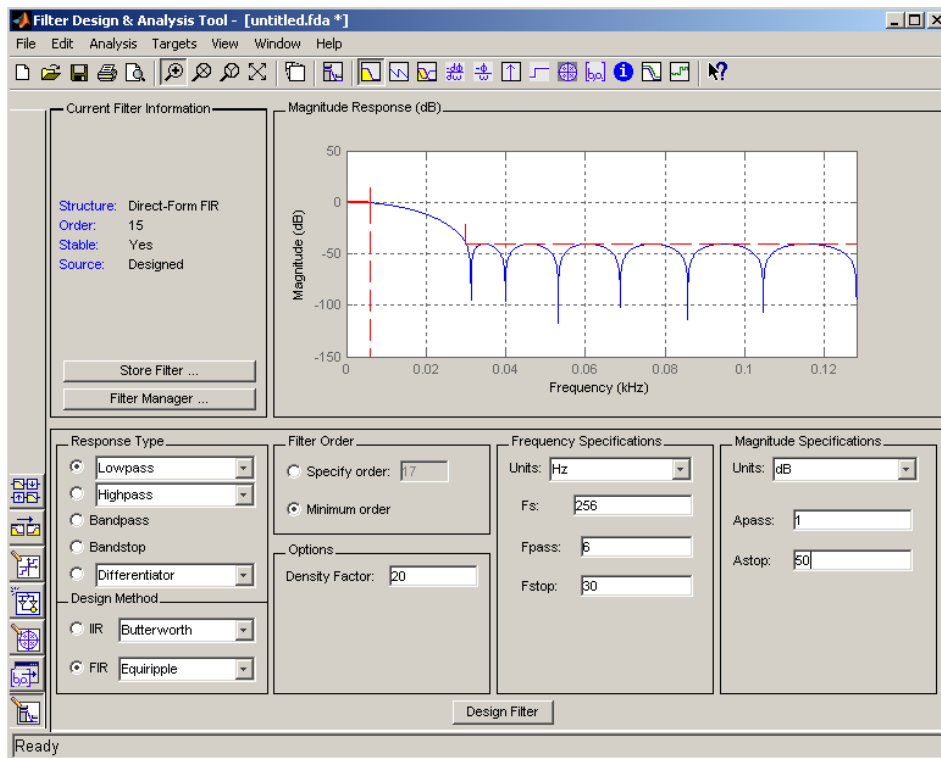


Figure 91 Illustration of Low Pass 17-tap Filter (without scaling) using MATLAB Filter Design Manager

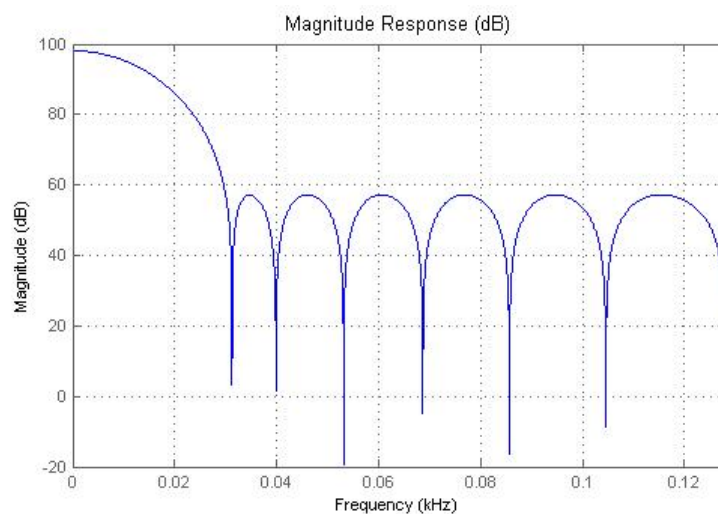


Figure 92 Plot of Magnitude versus Frequency Response for 17-Tap FIR Low-Pass Filter (gain factor of 500)

Similarly, using the filter design toolbox from MATLAB, the high-pass filter is designed with a corner frequency of 2 Hz as shown in Figure 93. The magnitude response of the 17-tap high-pass FIR direct form filter generated from MATLAB is shown in Figure 94.

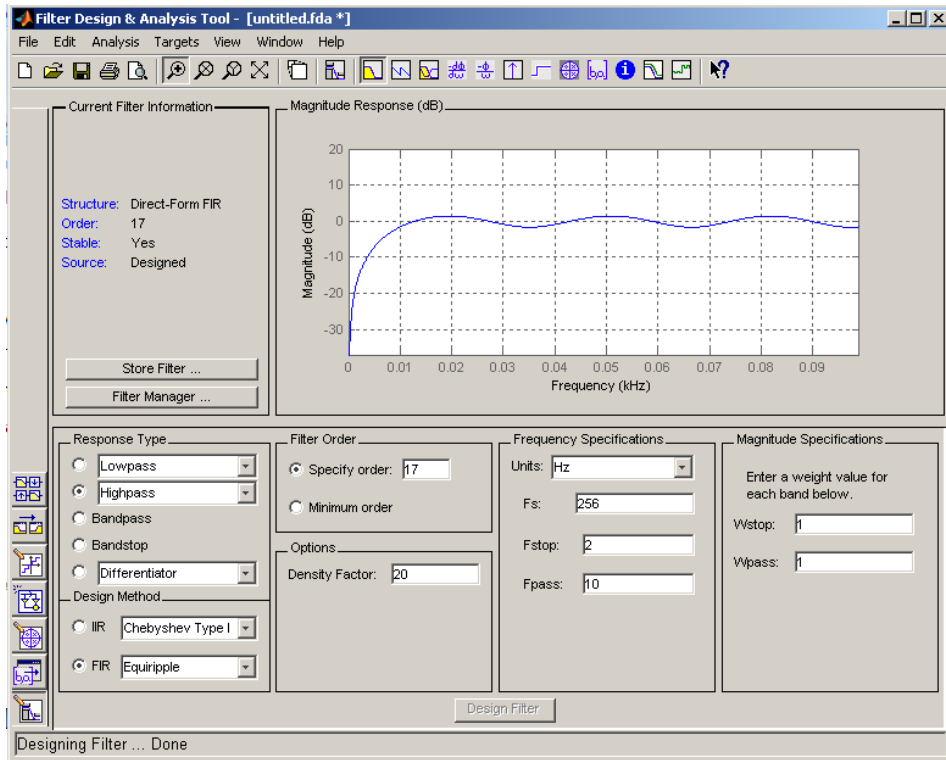


Figure 93 Illustration of High Pass 17-tap FIR Filter using MATLAB Filter Design Manager

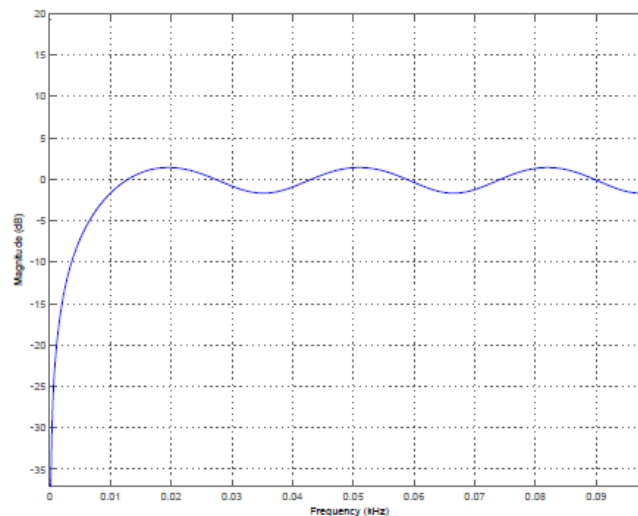


Figure 94 Plot of Magnitude versus Frequency Response for 17-Tap FIR High-Pass Filter

The filtered output is further processed by subtracting a fixed threshold from the filtered output. This cuts off the unwanted disturbances caused by the movement-related artifacts. Using this method, the peak of the MMSB signal can be discriminated from the complete MMSB waveform through the zero crossing detection. Using the model illustrated in Figure 89, the peak detection functionality is verified to work correctly through the waveforms as shown in Figure 95. In Figure 95, it can be observed that the peak of the simulated MMSB signal can accurately detected for each heart beat through the digital pulse output of the zero crossing detection block as shown in the bottom waveform.

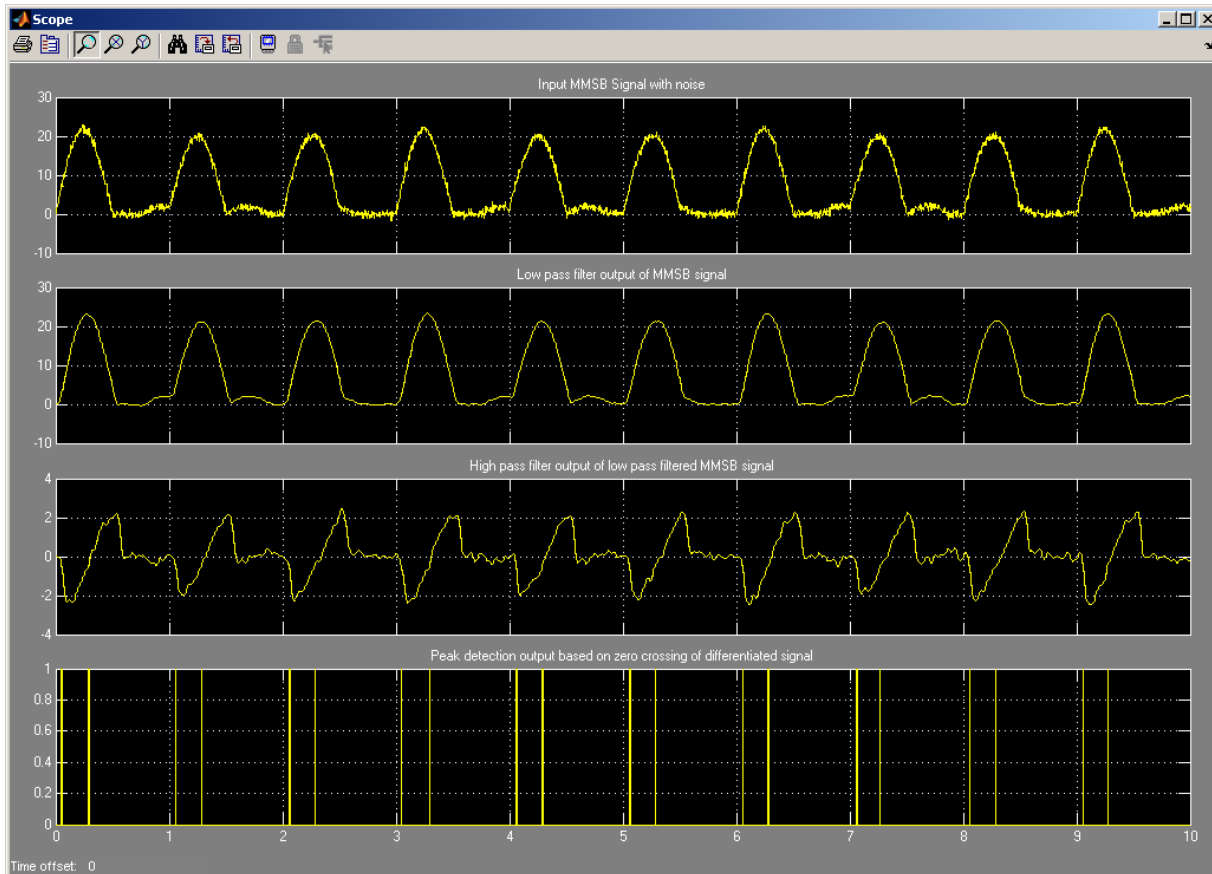


Figure 95 Output waveform of peak discriminator design

With the successful implementation of the peak detection architecture in Figure 89 using MATLAB / Simulink, the coefficients for the low pass and high pass filters are implemented in the hardware interface development software LabVIEW. In addition, in order to minimize error due to motion artifacts, the number of heart beats per minute is calculated using a three beat average. A C-function is implemented in LabVIEW where two variables in the C main function, counter and pulse_period, accurately track the time scale. Each output sample from the MMSB peak discriminator is compared against a set threshold to detect the presence of a beat. Pulse_period is incremented by one during every sample period. Because each sample occurs every 0.1 second, it is easy to track the time scale based on the number of counts in the pulse_period variable. A 4-sample time window is used as a debounce time using counter. Every time a beat is detected, counter is reset and an icon is turned on to represent the heartbeat. If a beat is not detected for 4 consecutive samples, a separation between successive beats is identified and the icon is turned off. The pulse_period is accumulated for three consecutive beats. On the third beat, pulse_period is used for the calculation of heart-rate per minute as shown in Equation 71 and reset for the next calculation.

$$\text{Equation 71} \quad \text{Heartbeat per minute} = \frac{1}{\left[\frac{\text{pulse_period}}{3 \times 10 \times 60} \right]} = \frac{1800}{\text{pulse_period}}$$

2.4 Packaging of MMSB blood pulse detection as a wearable device

In order to package sensor and magnet as a wrist wearable device to measure pulse rate without degradation of the MMSB sensitivity and system performance, a novel packaging method was developed and patented [9]. The novelty in the method of packaging relies on the use of air-bubble to achieve a constant air-gap separation between the magnet and the synthetic leader belt. This will allow the magnetic field coupling to be independent of the material used for the belt and at the same time, the belt will also provide physical conformity of blood vessel and the air-bubble as shown in Figure 96. In addition, the air-bubble will also minimize the pressure applied on the blood vessels resulting in a constriction of the blood vessel due to the wrist strip pressing against it.

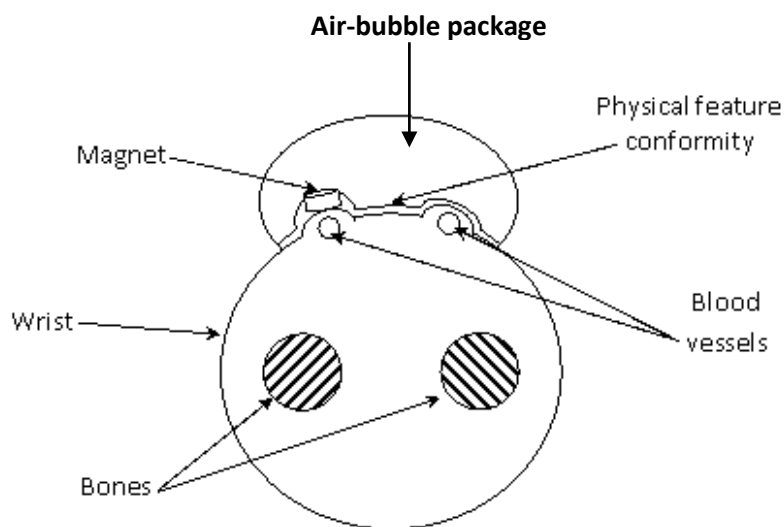


Figure 96 Illustration of packaging of MMSB blood pulse detection as a wearable device

Applying the method of packaging described in Figure 96, the final configuration for a wearable device applying the MMSB method of pulse rate measurements is shown in Figure 97.

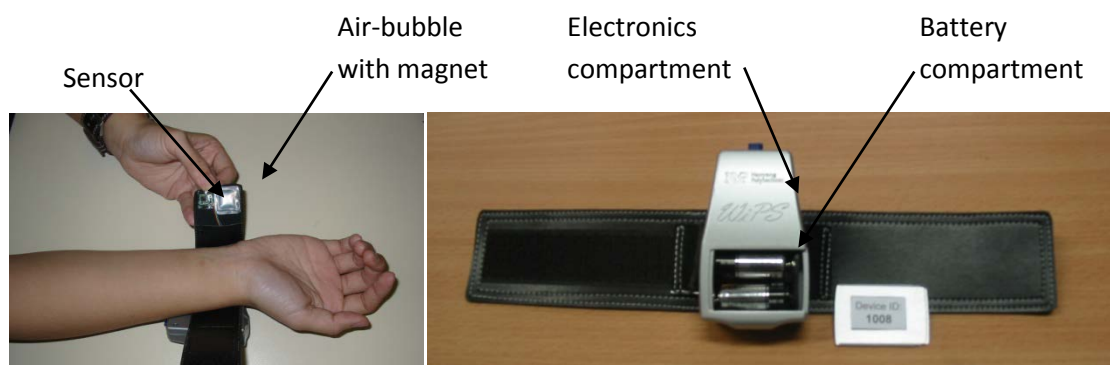


Figure 97 Final configuration of wearable device with wireless transmission

With the completion of the wearable device with wireless transmission, it was tested on 20 healthy users aged 20-25 and it was able to acquire their pulse rate where monitoring is achieved at the base

station. Using finger based pulse oximeter as the measurement reference, the pulse rate difference between the MMSB method and the pulse oximeter is tabulated in Table 22, where the maximum percentage difference is found to be less than 5%.

Subject	MMSB device	Pulse oximeter	Percentage difference
Subject 1	60	62	3.33%
Subject 2	78	77	-1.28%
Subject 3	68	70	2.94%
Subject 4	72	74	2.78%
Subject 5	89	91	2.25%
Subject 6	77	74	-3.90%
Subject 7	73	70	-4.11%
Subject 8	56	55	-1.79%
Subject 9	69	66	-4.35%
Subject 10	75	72	-4.00%
Subject 11	78	80	2.56%
Subject 12	82	85	3.66%
Subject 13	86	82	-4.65%
Subject 14	76	75	-1.32%
Subject 15	78	80	2.56%
Subject 16	69	72	4.35%
Subject 17	71	74	4.23%
Subject 18	79	80	1.27%
Subject 19	92	90	-2.17%
Subject 20	86	85	-1.16%

Table 22 Tabulation of heart rate measurements for MMSB device and Pulse Oximeter

In addition, this development was tested by the scientists in the medical research laboratory of the defense organization in Singapore and was found to be working well within the accepted tolerance [8].

To enhance this development for applications such as a personal lifestyle pulse rate monitoring device without wireless transmission, the algorithms developed in this section were implemented into an embedded signal processing system with an integrated LCD display. This will allow the PC based processing to be implemented into a standalone wearable pulse rate monitoring device.

2.5 Standalone pulse rate monitoring

The standalone pulse rate monitoring device was developed to allow monitoring of heart rate when the wireless network is not available. This will function as a lifestyle device for healthy subjects who are interested in self-monitoring of their heart rate continuously. As such, the lower cost INA321 amplifier (please refer to ANNEX B for datasheet of INA321) was selected in place of the INA118 so as to reduce the total cost. In addition, the INA321 is also specified to operate with a minimum of 2.7-V single rail power supply allowing the wearable device to operate based on a single cell Li-ion battery.

For this application, The INA321 is designed to provide a fixed amplification of 5 times for the MMSB signal instead of the 100x for the INA118. This is because of the availability of a high-gain amplifier in the signal processor which will be utilized in this design to provide a total of 2 stages of amplifications.

Table 23 shows the comparison of the technical specifications of INA321 with respect to the more expensive INA118. From Table 23, the INA321 has a specified CMRR of 90 dB, which is lower than that of INA118, is sufficient to reject the common-mode noise signals including the line frequency and its harmonics. However, the quiescent current of the INA321 is 40 μ A, and the shutdown mode current is less than 1 μ A allowing the amplifier circuit to operate with minimum power as compared to the INA118.

Parameters	INA118	INA321
CMRR minimum (dB)	107	90
Quiescent Current (+/-) (typ) (mA)	0.35	0.04
Approximate price (USD)	4.80 1ku	1.10 1ku
Shutdown current (mA)	Not available	< 0.01

Table 23 Comparison of critical specifications for INA118 and INA321

The signal processor used for the implementation of the embedded processing is the Texas Instrument MSP430FG439. The MSP430FG439 is a 16-Bit Ultra-Low-Power microcontroller unit (MCU) that comes with 60KB Flash, 2KB RAM, one 12-Bit Analog-to-Digital Convertor (ADC), two Digital-to-Analog Convertor (DAC), Direct memory access (DMA), integrated three operational amplifier (OPAMP) and interface control to support LCD display of 128 segment. Using one of the three integrated operational amplifiers in the Signal Processor, MSP430FG439, the final circuit configuration for the MMSB amplification circuit is shown in Figure 98, which includes the INA321, the OAO, the analog integrator OA1 and digital-to-analog converters DAC1. Through INA321 and OAO,

the MMSB signal will be amplified with a total gain of 500. As the gain is very high, slight imbalances in the sensor differential outputs will cause the common-mode signal to offset, resulting in noise at the OA00 output. In addition, the output will also be sensitive to the variations in positional alignment between blood vessel and sensor. This will result in a variation of the dc content of the amplified differential signal and manifests itself as a drift in the baseline of the MMSB, commonly known as baseline wandering.

To manage the baseline wandering, an analog integrator scheme is designed with the use of OA1, the second of the three integrated operational amplifiers in the Signal Processor, MSP430FG439 (reference Figure 98). The integrator integrates the DC content of the 5 times amplified MMSB and feeds it back to the INA321. The feedback allows the INA321 to maintain a constant DC level at the output.

In addition, the digital baseline wandering scheme is implemented through the use of the DAC1, one of the two integrated DAC12 12-bit digital-to-analog converters in the Signal Processor, MSP430FG439 (reference Figure 98). Using DAC1 allows the user to implement a digital baseline wandering scheme through software implementation of digital integration.

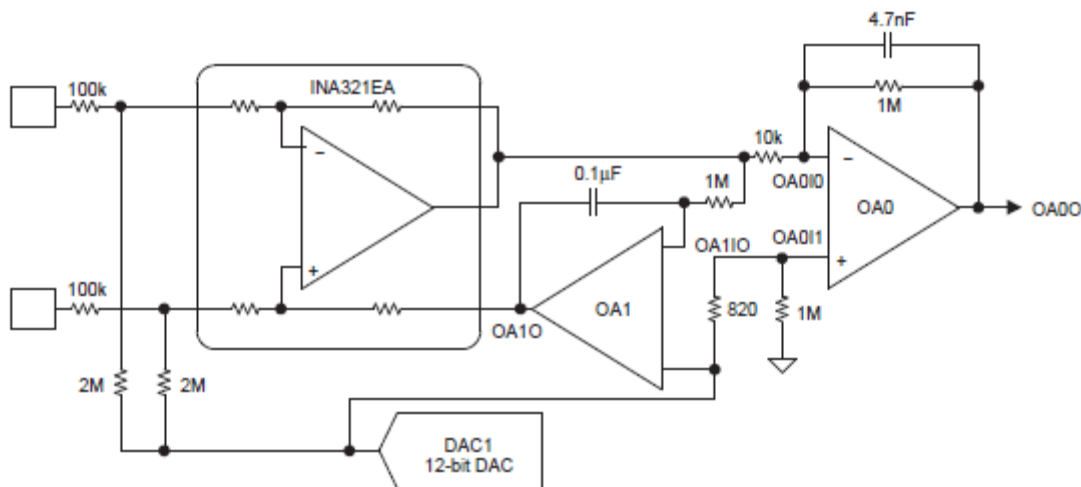


Figure 98 MMSB amplifier circuit and voltage reference feedback design

The final block diagram of the standalone pulse rate monitoring design is shown in Figure 99 where the final schematic and PCB of size 3.5cm x 3cm are shown in Figure 100 and Figure 101 respectively. Using the algorithm developed and tested in the LABVIEW environment (described in section 2.3 of this Chapter), the C-code is re-targeted for implementation on the Signal Processor, MSP430FG439.

The pulse rate was measured for 20 healthy subjects age 20-25 using the MMSB embedded device and a commercially available pulse oximeter as tabulated in Table 24. Results from this comparison

show both devices measures similar heart rate for all subjects with a maximum difference of less than 5%.

Subject	MMSB Embedded device	Pulse oximeter	Percentage difference
Subject 1	61	63	3.28%
Subject 2	75	77	2.67%
Subject 3	66	68	3.03%
Subject 4	75	74	-1.33%
Subject 5	91	91	0.00%
Subject 6	75	76	1.33%
Subject 7	73	71	-2.74%
Subject 8	60	58	-3.33%
Subject 9	70	66	-5.71%
Subject 10	75	74	-1.33%
Subject 11	78	80	2.56%
Subject 12	80	82	2.50%
Subject 13	90	88	-2.22%
Subject 14	76	75	-1.32%
Subject 15	75	76	1.33%
Subject 16	67	70	4.48%
Subject 17	71	74	4.23%
Subject 18	76	73	-3.95%
Subject 19	93	90	-3.23%
Subject 20	80	82	2.50%

Table 24 Tabulation of heart rate measurements for MMSB Embedded device and Pulse Oximeter

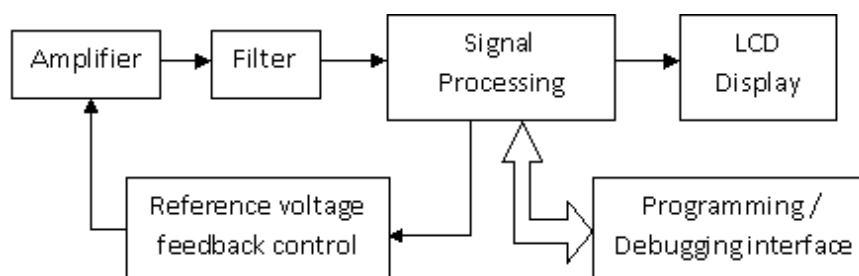


Figure 99 Block diagram of standalone pulse rate monitoring design

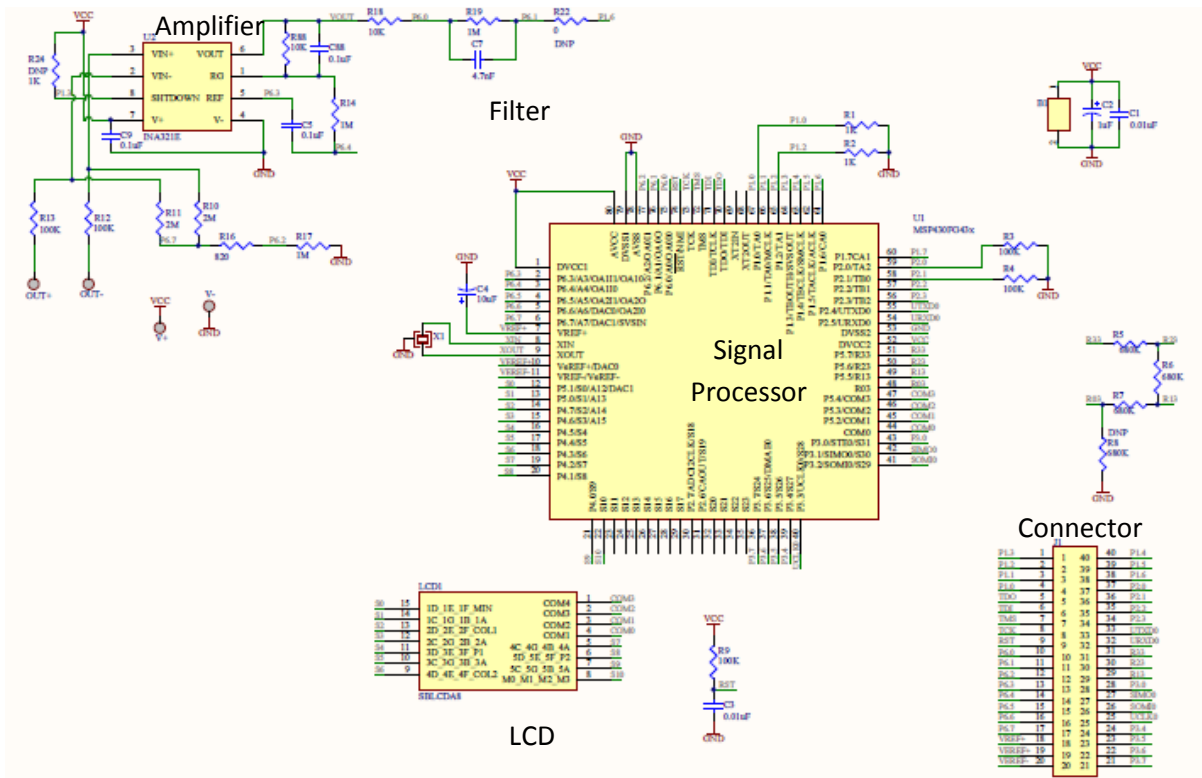


Figure 100 Schematic design of the standalone wearable pulse rate monitoring

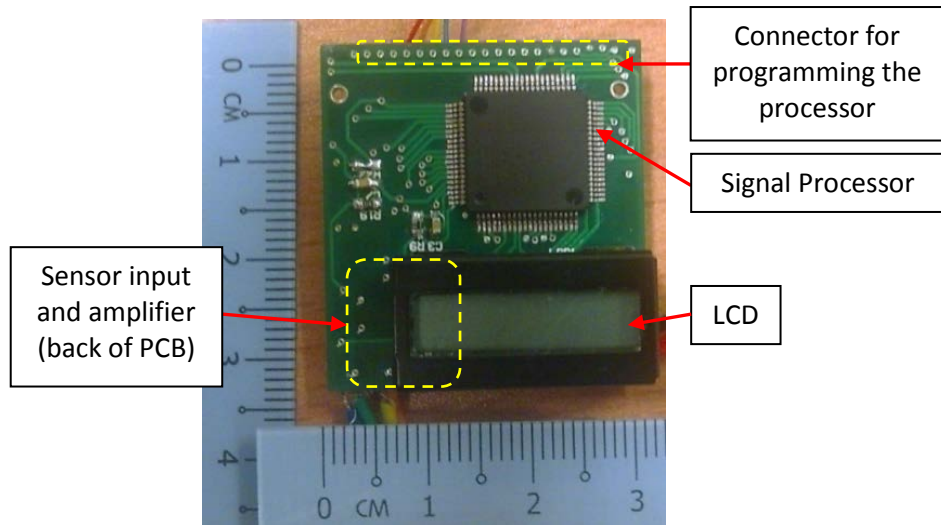


Figure 101 PCB design for standalone wearable pulse rate monitoring

2.6 Conclusions

With the successful implementation of the wearable pulse rate monitoring device with either wireless or local LCD display of results, the algorithm developed is proven to be able to acquire the heart rate using MMSB signal. In addition, the successful comparison of the measured pulse rate with commercially available pulse plethysmogram demonstrates the feasibility of the application of MMSB as a lifestyle device.

In the next section, the development will be focused on the application of MMSB to acquire blood volume pulse to assess the mental state of being relaxed or stressed of human subjects. This development is in collaboration with a commercial company where the outcome will be implemented into a commercial product.

3 Relaxation assessment using blood volume pulse

It has been reported in various publications that there are many methods available in medical research to determine the mental state of being relaxed or stressed of a human being. Studies using various physiological vital signs such as the body thermal state [1], blood perfusion and oxygen near the brain [3], brain electrical activities [4] and many others, had been conducted to understand its correlation to the mental state of the human subject. Each of these measurements of physiological vital signs has its advantages and limitations as these are measurements of the outcome of the mental state of the subject. As such, the accepted method in the medical research community to access the mental state of the brain for human subjects is through the measurement of the brain's electrical activities, electroencephalography (EEG). However, the method of measuring EEG is found to be inappropriate for commercial applications as it involves a series of procedure such as wearing of the EEG cap, skin preparation, electrolyte application, EEG signal calibration and substantial amount of signal processing. Therefore, various studies had been conducted to assess the mental state of the user through the measurements of physiological vital signs as the surrogate markers.

In order to understand the frequency of using various non-invasive methods to measure physiological vital signs so as to assess the mental state of being relaxed or stressed of the subject, a review, based on 20 publications, was conducted. The results of this review was summarized in Figure 102 and it was found that Blood Volume Pulse (BVP) is the most frequently quoted method (33.33%) to assess the mental state of the subject (details of this study is documented in ANNEX B). This is followed by heart rate variability (30.77%) and skin temperature (12.82%). As the MMSB signal contains both the BVP and pulse^{****} rate variability, it is potentially feasible to use MMSB to assess the mental state of being relaxed or stressed for subjects under observation. For the development work outlined in the following section, the BVP is selected as the measurement to assess the mental state of being relaxed or stressed. This is because the stimulus chosen for this work is a massage chair and it has been reported [5] that an increase of blood flow was found for subjects who enjoy being massaged while a decrease in blood flow was found for subjects who do not like being massaged. Such a measurement will therefore support the assessment of the physical state of being relaxed or stressed for subjects after a massage.

**** Pulse rate variability has been reported to be an acceptable alternative to heart rate variability when ECG recording is not available [10].

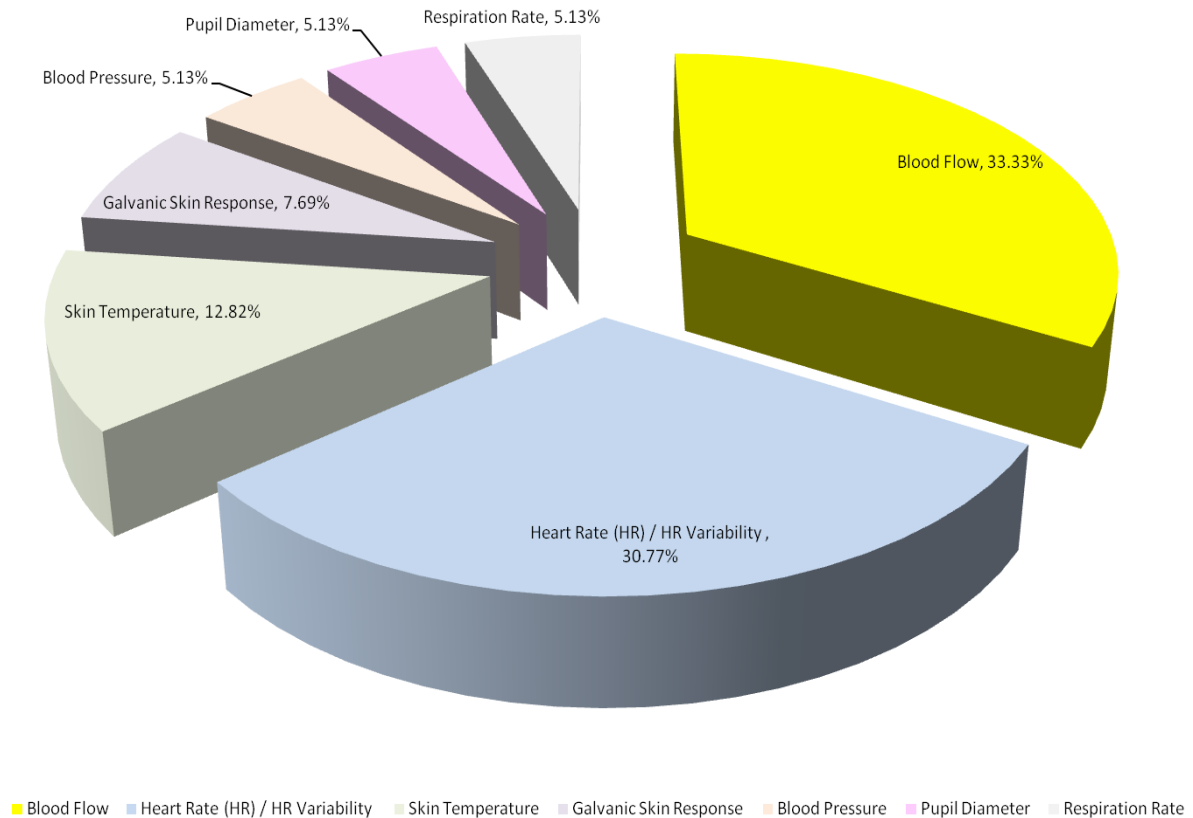


Figure 102 Summary on the frequency of non-invasive methods used to assess the mental state

3.1 Introduction

Blood volume pulse is a psycho physiological index showing the activation of the autonomic nervous system. Its simple recording, simple waveform and skin application make this index appealing for psycho physiological studies. Investigation of the direct effects of neck massages using blood volume pulse was performed in [5] and concluded that short-term massage on the neck has a direct influence on blood volume pulse and indeed tends to increase its value. In addition, one of the important observations concluded in this experiment is the result of neck massage bringing about a general relaxation to the subjects.

In a separate study conducted at Florida International University, four signals: Blood Volume Pulse (BVP), Galvanic Skin Response (GSR), Pupil Diameter (PD) and Skin Temperature (ST) were concurrently acquired and analyzed to differentiate affective states in the user [6]. In this experiment, several signal processing techniques are applied to the signals collected to extract the most relevant features in the physiological responses and feed them into learning systems, to accomplish the affective state classification. Three learning algorithms are applied to this classification process and their performance is compared. Results in this experiment indicate that

the physiological signals monitored do, in fact, have a strong correlation with the changes in emotional state of the experimental subjects.

Finally, a medical research was conducted [7] on patients with coronary artery disease (CAD) to determine if mental stress may provoke ischemic electrocardiograph changes and abnormalities in regional and global left ventricular function. This study stress the both healthy and CAD patients using mathematic subtractions in a progressively challenging sequence and concluded that CAD patients exhibit an attenuated blood flow response to mental stress that may contribute to mental stress induced ischemic episodes in daily life.

For both of the above experiments conducted in United State [6][7], it was documented that the Laser Doppler Blood perfusion measurement was used. Therefore, this section of development work will outline the study and implementation conducted using the OSIM iDesire Massage chair as the relaxation stimulus where the change in blood volume pulse was concurrently assessed using MMSB based sensing system embedded in the OSIM iDesire Massage Chair and the Laser Doppler Blood perfusion equipment. The change in blood volume pulse was correlated to the relaxation state of the muscle providing the provision to infer the mental state (e.g. relax or stress) of the subject.

3.2 Experimental overview

The design of a system to assess the mental state of being relaxed or stressed of a human subject has to be able to acquire the MMSB signal with minimum interference to the relaxation or stress stimulus and minimum discomfort to the subject. As such, the system design for this section will first explore the optimal locations to mount the sensor and magnet in order to acquire MMSB on the massage chair. This is followed by the verifications of the reliability of the measured signal through measurements of the MMSB signal on 5 subjects with different demography. With the reliability measured, the location with the best reliability will be selected and signal processing will be implemented to remove the noise from the massage chair. Finally, the experiment stimulus for relaxation will be selected and repeated measurements will be done on each of the 5 subjects to ensure repeatability of experiment. Finally, a statistical approach will be developed to determine the change in BVP. The BVP obtained using the statistical analysis will also be compared with measurements using the medical research instrument that applies the laser Doppler method to measure blood volume flow. Once the system design is verified to be reliable and repeatable, a trial, consisting 40 subjects, was conducted using the developed system to assess the mental state of being relaxed or stressed after the application of the massage as the stimulus.

3.2.1 System setup for trial

The MMSB signal characteristics measured on the heel is similar to that obtained on the wrist. As such, the amplifier circuit designed in Figure 86 was used for this experiment and trial. The output of the amplifier circuit is directly connected to the data acquisition card from National Instrument where pre and post massage MMSB signals are collected and analyzed using a personal computer as shown in Figure 103.

The software LabView is used on the personal computer to process the measured data for change in BVP before and after massage. Using the relationship between amplitude and BVP described in Chapter 4, the amplitude of each pulse is measured to indicate the relative change in BVP value before and after a massage. From the MMSB waveform recorded, the peak amplitude of each pulse is manually extracted and consolidated for statistical processing in the section 3.4.

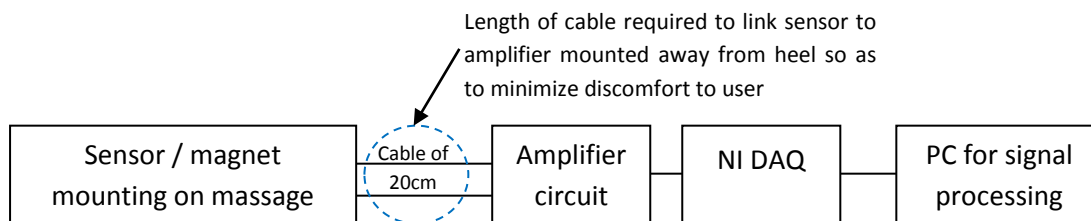


Figure 103 Illustration of the circuit configurations for BVP analysis on massage chair

3.2.2 Mounting of sensor on the massage chair

In order to achieve MMSB based blood volume pulse measurement on the subject with minimum discomfort, various locations on the OSIM iDesire massage chair were explored. As the back massage function on the massage chair is based on mechanical rollers, it is not desirable to consider the upper part of the back body for mounting of sensor as the rollers may cause damage to the sensor.

Using the lower body as the considerations, a few potential locations for MMSB blood volume pulse measurement are identified and illustrated as shown in Figure 104.

Based on the ease of assessing the major blood vessels on the skin and the presence of air-bags on the OSIM iDesire Massage chair, a total of 6 locations were identified. These are the calf, kneel (underside), thigh, tail-bone region, wrist and heel as illustrated in Figure 104.



Figure 104 Illustration on the potential locations to acquire MMSB signal on the massage chair

In order to select the location which will yield the most consistent MMSB results for all subjects, a series of experiments were carried out to determine the most appropriate location for MMSB signal acquisition. In these experiments, sensors and magnets were mounted on each of these locations (as shown in Figure 105) and MMSB signals were acquired before and after the massage. These experiments were repeated on 5 subjects of different demography and aged. Through these experiments, it was found that the underside of knee and the heel were the only locations that will produce consistent MMSB results on the massage chair. The rest of the locations were found to have challenges such as the inconsistency of air-bags to hold the calf with consistent grip for all subjects; non-conformance of air-bags on the wrist and thickness of clothing at the tail-bone region.

In order to implement MMSB signal acquisition on the underside of knee, it will require the addition of airbags to the massage chair (refer Figure 105) and was found to be inappropriate for commercialization due to the cost and user comfort.

As such, the heel was selected as the location that is most appropriate for MMSB signal acquisition on the OSIM iDesire massage chair. This is because the iDesire Massage chair is equipped with airbag at the heel and includes heel massaging function which will inflate the airbag at the heel to provide the necessary grip and support to acquire MMSB signal reliably. In addition, the location of the blood vessel on the heel was found to exhibit minimum differences for most of the subjects with feet size ranging from 5 to 12 (European size).

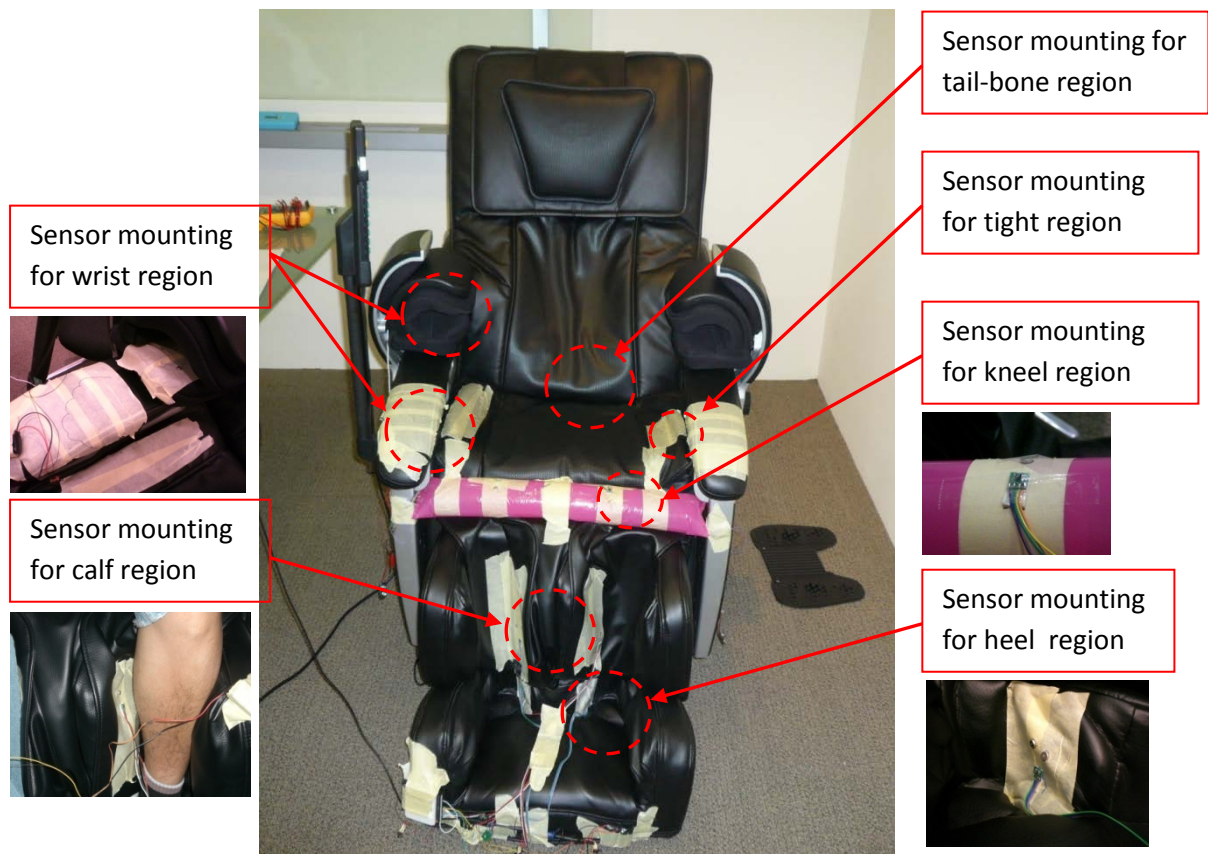


Figure 105 Illustration on mounting of sensors and magnet on OSIM massage chair

In order to ensure reliable signal acquisition on the heel for subjects with feet size ranging from 5 to 12, a series of experiments were conducted and it was found that the configuration of two magnets and a single sensor (as shown in Figure 106^{§§§§§}) at the simulated optimal distance of 2cm apart (Reference Chapter 3), as shown in Figure 107, provides the most consistent results. Using this configuration, the sensor and magnet mounting was able to acquire the MMSB signal reliably during the inflation of the airbags.

In addition, for the purpose of commercialization, the ability of MMSB to measure through clothing was also found to have an added advantage as the user do not need to take off their socks during massage. Such an advantage will provide ease of usage for the user as the system will be able to assess the mental state of being relaxed or stressed independent on the user's choice of clothing.

^{§§§§§} Due to the fact that this design will be commercialised, the final configuration of the sensors and magnets array is not shown.



Figure 106 Illustration of sensor mounting on OSIM iDesire Massage Chair

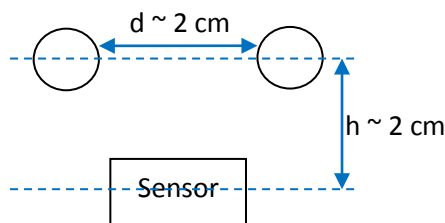


Figure 107 Illustration of sensor and magnets position

3.2.3 Selection of a stimulus for relaxation assessment

The OSIM iDesire Massage chair has eight pre-defined massage sequences and has also included the options to manually overwrite any of the pre-defined massage sequence to achieve user desired combinations. In order to ensure repeatability for the measurement on the change of BVP before and after a stimulus, the pre-defined massage function labelled full-body relief was selected as the stimulus for relaxation assessment.

With the selection of the massage stimulus, heel massage will also need to be included before and after the stimulus to acquire BVP using MMSB signals. This is because the heel massage will include a series of sequence where the airbags at the heel are inflated to allow the sensor and magnet to reach the heel of the subject thereby allowing MMSB signal to be acquired.

3.2.4 Filtering of EM noise and DC drift

During the experiments, it was found that the massage chair generates a significant amount of electromagnetic noise due to the switching of the various integrated relays. The noises generated

will affect the quality of the MMSB signal as these are impulsive and typically large in amplitude. In addition, due to the inflation of the airbags, it was also found that the DC level drift is continuously changing as the sensor and magnet are moving on the airbag. These problems can be observed using the data acquisition modules from National Instrument where a typical signal is shown in Figure 108. It can be observed that the noise is impulsive and the DC drift is dominant. As such, without any signal processing, the MMSB signal cannot be easily identified from Figure 108.

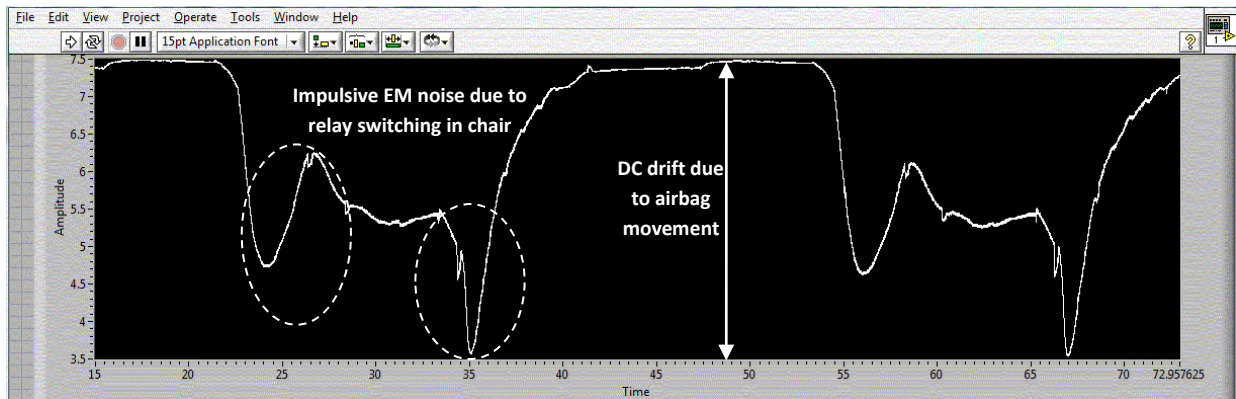


Figure 108 Typical signal acquired on by MMSB sensor on OSIM iDesire Massage Chair

In order to remove the electromagnetic noise and DC drift, an Infinite Impulse Response (IIR) bandpass filter was designed in LabVIEW to remove the DC drift and the electromagnetic noise. The IIR bandpass filter was selected due to its inherent characteristics of DC drift removal through autoregressive-moving average (ARMA) filtering, operating on current and past input values and current and past output values. The inherent characteristics can be observed through the architecture of the IIR filter as shown in Figure 109, where z^{-1} block represents a unit delay and the coefficients and number of feedback or feedforward paths are determines the order to the IIR filter.

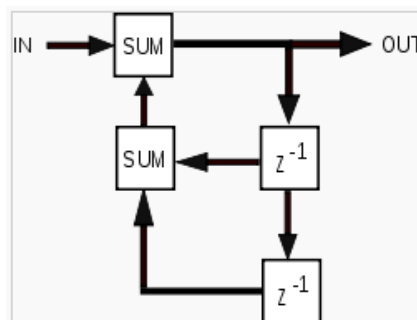


Figure 109 Architecture of a simple 2nd order IIR filter

In addition, IIR filters can achieve much sharper transition region roll-off than FIR filter of the same order. This is important for the IIR filter designed in the LabView environment as it will be implemented in an embedded signal processing system with minimum resources (i.e. low order IIR filter) and high selectivity of signal.

Therefore, the characteristics of the 2nd order IIR Filter implemented in this design is shown in Figure 110, where the 3dB passband is designed for 0.5 to 7 Hz with fast roll off to remove out-of band-signals with minimum attenuation to the signal strength. The choice passband to start from 0.5 Hz is to remove the DC drift, which will result in the baseline wondering of the signal. The 3dB cut-off frequency of 7 Hz is selected the typical heart rate of human being 1-1.5 Hz. In addition, the high selectivity obtained from the filter is designed to cater to the high electromagnetic interference due to the motorized movement of the chair.

The topology for the IIR filter is selected based on maximally flat magnitude filter so that the frequency response is as flat as mathematically possible in the passband. In this way, the frequency characteristics of the MMSB signal will not be affected.

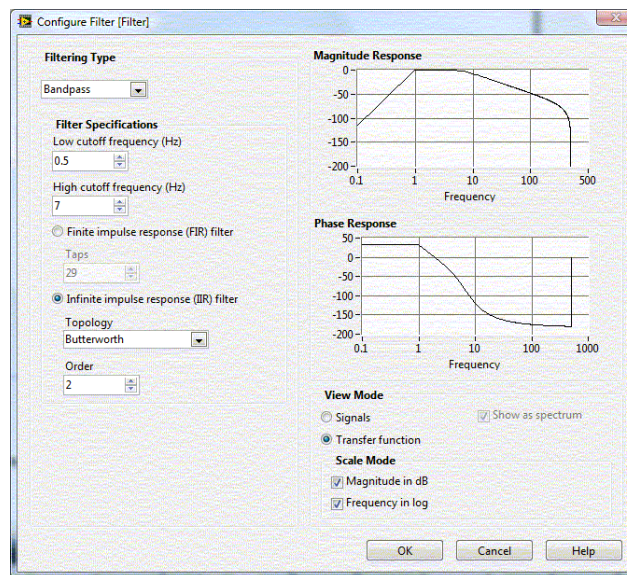


Figure 110 IIR filter design using LabVIEW

Using the designed parameters in Figure 110, it can be observed that the magnitude of the passband is relatively flat due to selection of the Butterworth topology for the IIR filter. The Butterworth topology has the advantages of having smooth, flat and monotonically decreasing frequency response allowing the MMSB signal to be selected with minimum distortion.

With the performance of the IIR filter simulated and implemented in LabVIEW, the filter is integrated into the signal processing block as shown in Figure 111. The typical unfiltered signal shown in Figure 108 was processed through the filter and the output is shown in Figure 112. Based on results obtained in Figure 112, it is found the IIR filter is able to remove most of the DC drift. This will result in the ease of MMSB signal identification after filtering. To study the effectiveness of the filter in removing the noise and DC drift, the waveform is plotted at a higher resolution (or smaller time step)

as shown in Figure 113, where it can be observed that the DC drift was removed completely and the impact of the impulsive EM noise on the signal quality is minimized.

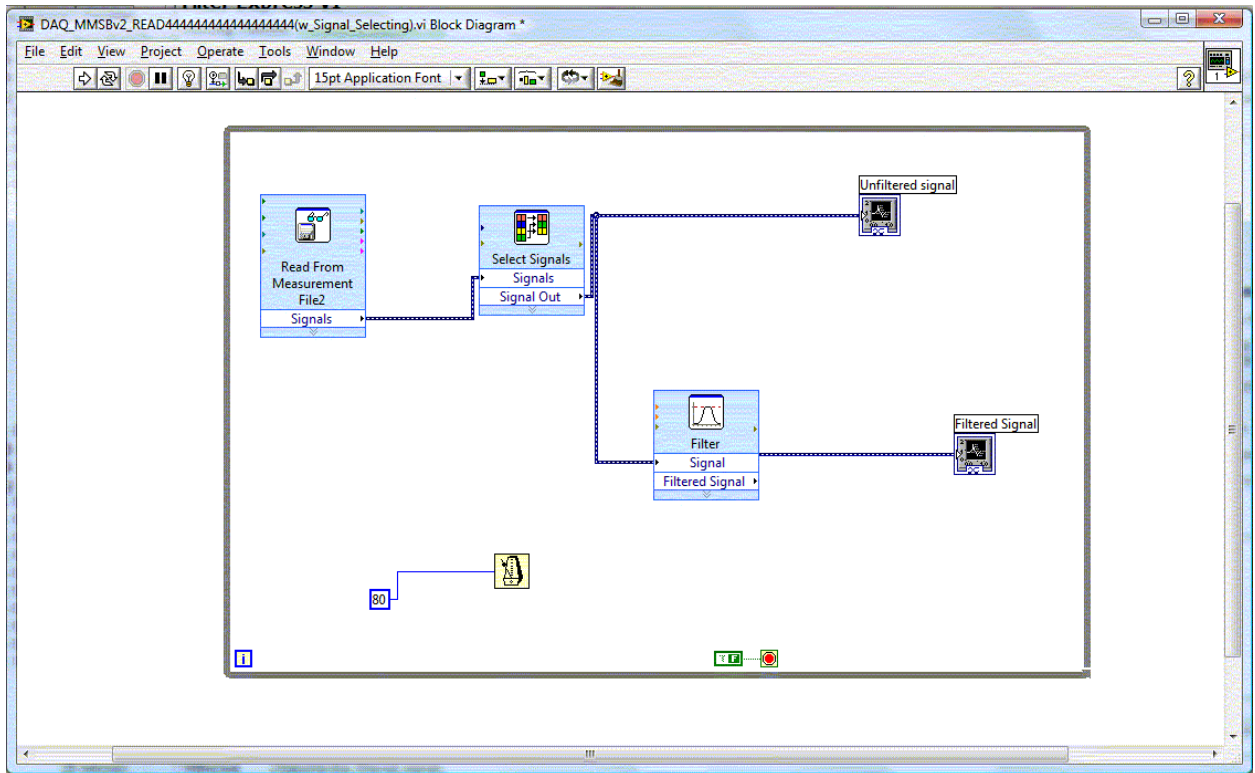


Figure 111 Implementation of the IIR filter for MMSB signal filtering

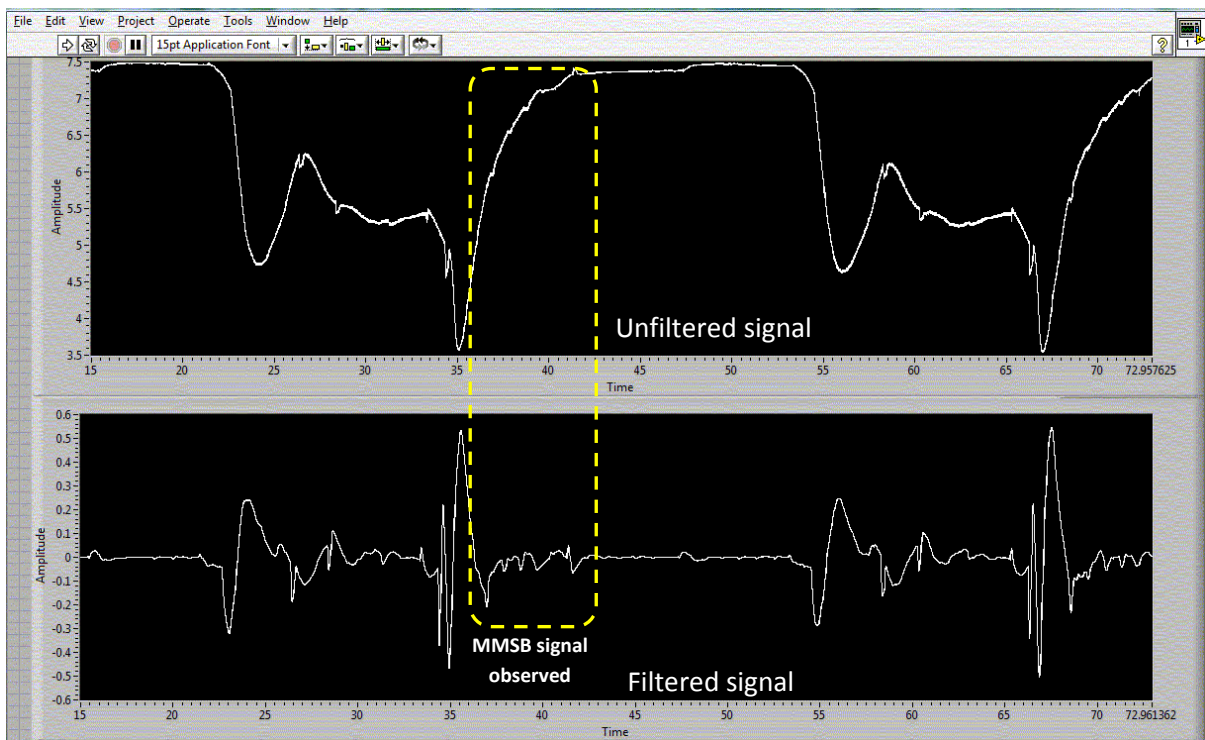


Figure 112 Typical signal output comparison between filtered and unfiltered signal

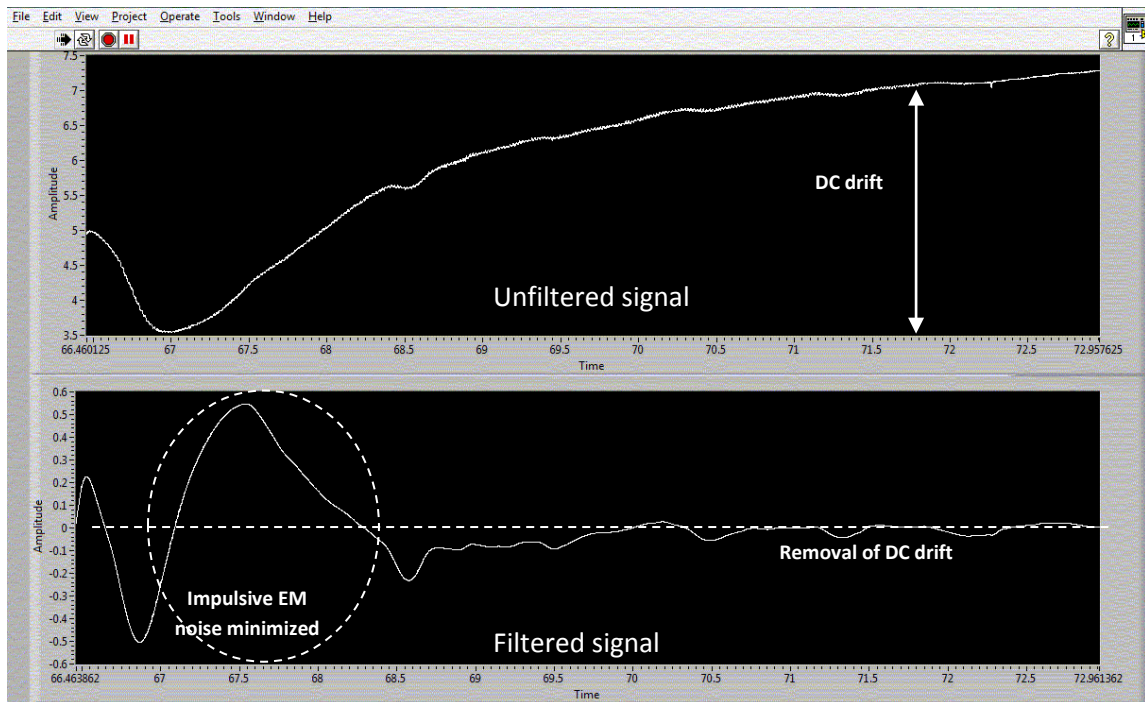


Figure 113 Typical response of the IIR filter in removing DC drift and impulsive EM noise

3.3 Setup and procedure used in trial

Using the typical heart rate of a healthy adult as 1 to 1.3Hz (i.e. 60 to 80 pulses/minute), the heel massage will have to acquire MMSB signal for at least 5 minutes so as to ensure that there is enough BVP pulses available for analysis. This is because the heel massage is based on the pre-programmed sequence which will inflate the airbag at the heel in intervals of twice per minute and each grip will typically yield 3 to 5 detectable pulses for BVP analysis. Therefore, a total of 30 to 50 pulses will be collected during the heel massage for the statistical analysis of the BVP to assess the change in BVP before and after the full-body relief massage.

In addition, as the blood volume pulse is dependent on the position of the subject on the massage chair, the massage chair was adjusted to a pre-defined inclination (of approximately 60 degrees) as shown in Figure 114 for all subjects, after they are comfortably seated on the massage chair. With the subject at the recommended inclination angle, the heel and calf will also be well rested on the massage chair, which will be supporting most of the weight on the lower body. Thus, this setup will reduce error

in BVP measurements that can be introduced during the trial due to muscle tension related to the sitting position of the subject.



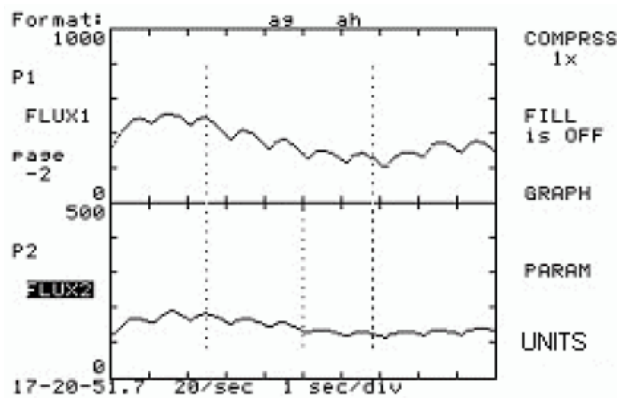
Figure 114 Inclination of OSIM massage chair during trial

In addition, the MMSB measurement is also compared with the laser Doppler measurements using the medical research gold standard, Moors Instruments. The objective of this comparison is to ensure that MMSB measurement is calibrated with the medical research gold standard.

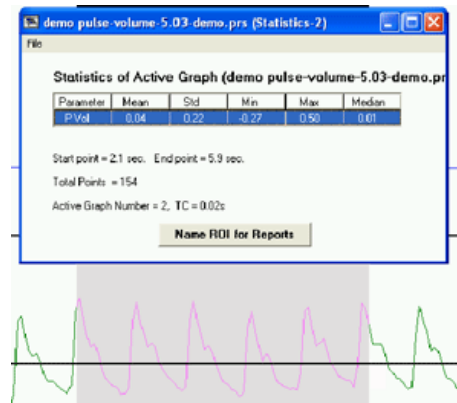
Typical mountings of the laser probes are shown in Figure 115, where the massage chair will not damage the probes during the experiment. With this mounting, the typical measured result using the blood laser Doppler instrument is shown in Figure 116.



Figure 115 Illustration of mounting for laser probe during trial



(a) Typical waveform obtained



(b) Waveform selection with statistical

analysis

Figure 116 Typical LDF signal acquired using Moors Instrument

Using the setup described above, the procedure for the experimental process to collect BVP data during the experiment and subsequent trials is outlined below.

1. Massage sequence on the heel to acquire blood flow information using MMSB (5 minutes)
2. Measurement of BVP using Moor's instrument (refer to Figure 115)
3. Full-body massage sequence (15 minutes)
4. Massage sequence on the heel to acquire blood flow information using MMSB(5 minutes)
5. Measurement of BVP using Moor's instrument(refer to Figure 115)

Data collection for MMSB and Laser Doppler are done sequentially. This is because the air-bag will have to be activated to achieve conformance grip of the heel so as to acquire MMSB data while the Laser Doppler measurements is highly sensitive to motion artefacts due to the heel massage and will have to be acquired when there is no massage activity.

Data collection for each subject participating in this trial was repeated twice so that the data collected are consistent. For each subject, a time gap of 3 hours is ensured between two measurements so that the effect of massage during the first experiment is minimized. In addition, the data collected will also be statistically independent.

Using the setup and the method for data collection as described above, the experiment was repeated on 40 subjects and the results are captured and analysed in LabView environment. The following sections will present the results obtained based on a statistical analysis of the measured BVP.

3.4 Data extraction for MMSB signals

For each of the data collected, the peak voltages, V_1 , V_2 and V_3 for each measured pulse is extracted in LabView environment using manual measurement of the peak to valley as shown in Figure 117.

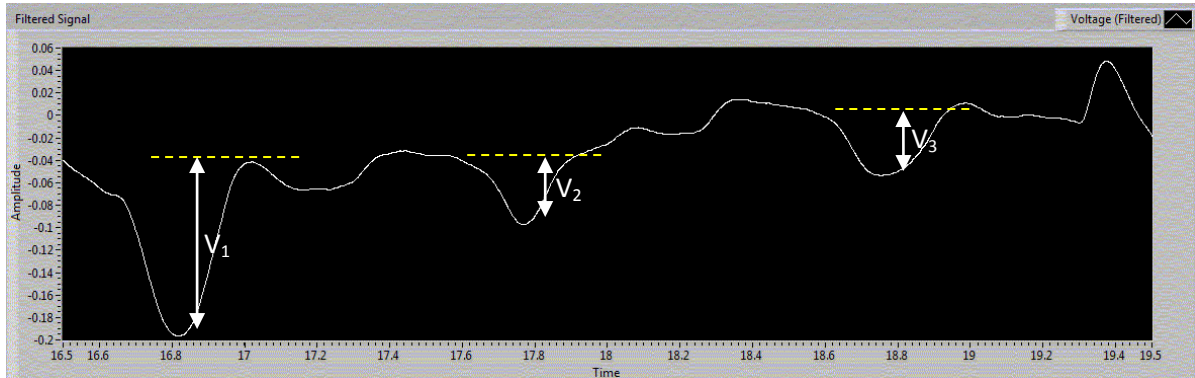


Figure 117 Illustration of peak to valley measurements of MMSB data for BVP data extraction

Using the peak to valley method of measurement, a typical set of data using MMSB signal acquisition for each experimental subject is extracted as shown in Table 25. For each experiment, 48 pulses are to be collected before and after the “full-body relief” massage sequence. This is to ensure that the data are spread over a period of time where each data is assumed to be statistically independent.

Time Step	Data points acquired during approx 5 min massage (mV)												Average of the Peak Voltages (mV)
	1st Peak	2nd Peak	3rd Peak	4th Peak	5th Peak	6th Peak	7th Peak	8th Peak	9th Peak	10th Peak	11th Peak	12th Peak	
1	30	50	40	40	20	20	80	70	70	80	40	30	48
2	30	50	40	35	30	30	80	80	70	70	60	30	50
3	50	60	40	30	40	30	90	80	75	60	55	40	54
4	60	40	30	40	40	40	80	80	70	70	50	30	53
5	50	50	40	40	50	40	90	100	90	80	80	50	63
6	60	70	50	50	50	40	80	100	100	80	70	40	66
7	75	60	40	50	50	45	80	110	90	80	70	50	67
8	70	70	40	40	40	40	85	100	100	70	60	50	64

Table 25 Typical data collected based on each trial

Similar data collection using the Moor’s instrument for BVP is also collected and each of the data set is analyzed statistically. In order to minimize errors introduced to the BVP results from Moors Instrument due to voltage drift of the optical sensor, the data used for the statistical analysis is typically taken along the plateau of the signal. In addition, the data acquired from Moor’s Instrument is processed based on the sampled data where each measurements used (around 10 seconds) contains at least 1000 points. This is done through the use of the Moor’s software that is able to provide statistical data based on a selected region of interest. As the data points used for Moor’s Instrument is typically large, the Z-statistical test will be used.

A Z-statistical test is the statistical test that is using the distribution of the test statistic under the null hypothesis where the data is assumed to approximate to a normal distribution. As the data from Moor’s instrument and MMSB both contains a relatively large sample (> 30 data points each), the Z-statistical test will be applicable. In this test, the means between two samples will be analysed using Equation 72.

Equation 72

$$Z = \frac{X_2 - X_1}{\sqrt{\frac{Std_2^2}{N_2} + \frac{Std_1^2}{N_1}}}$$

where X_2, X_1 = means of data before and after massage respectively

Std_2, Std_1 = standard deviation of data before and after massage respectively

N_2, N_1 = sample size of data collected before and after massage respectively

3.5 Statistical analysis of trial

The statistical data analysed from the BVP measured using MMSB is compared with the statistical data obtained using the laser Doppler medical instrument to measure BVP. A typical set of data obtained for this trial is shown in Figure 118. With the statistical data tabulated for both the Moor Instrument and MMSB, the two-tailed Z-test is used as inference to determine if the difference between the statistical values obtained for each of the measurement is either sufficiently small or sufficiently large.

Moor	Before						
	Parameters	Mean	Std	Min	Max	Median	N
	F1	3.07	0.33	2.33	4.01	3.05	1691
MMSB	Before						
	Parameters	Mean	Std	Min	Max	Median	N
	F1	26.8125	4.936431	24	29.00	25.00	48
Moor	After						
	Parameters	Mean	Std	Min	Max	Median	N
	F1	4.03	0.48	3	5.47	3.98	1492
MMSB	After						
	Parameters	Mean	Std	Min	Max	Median	N
	F1	34.45833	9.624787	28	39	35	48

Figure 118 Typical statistical data obtained before and after massage

The results obtained for the two-tailed Z-test at 95% and 99% confidence interval (CI) for both the Moors Instrument and MMSB are tabulated in Table 26. Based on the results obtained, it can be observed that the MMSB and Moors data have a 93.3% correlation (i.e. similar) at 95% confidence

interval. However, at 99% confidence interval, the data was found to have 91.3% correlation (i.e. similar).

The two-tailed Z-test at 99% CI produces a lower correlation due to the fact that the test is more stringent at 99% CI as compared to 95% CI.

In addition, the results obtained were also analyzed for the average number of cases where the subject's BVP improves after massage through the use of the Z-test. For cases where a positive significant was found, it can be concluded that the subject's BVP improves significantly after the massage. Therefore, using MMSB as the method of BVP measurements, at both 95% and 99% CI, the average number of measured cases with improved BVP is 61.25%. Comparing this to the 63% percent of cases reported to have improved BVP after massage [5], the results are concluded to be comparable.

	Moors@95% CI	MMSB@95% CI	Moors@99% CI	MMSB@99% CI
Positive Significant	30 cases	28 cases	23 cases	21 cases
Negative or No Significant	10 cases	12 cases	17 cases	19 cases

Table 26 Results obtained for two-tailed Z-test using MMSB and Moors Instrument

3.6 Conclusions

The results obtained in this study were found to be repeatable and the MMSB data correlates well with the measurements using laser Doppler method (Table 26). In addition, the number of cases with improved BVP correlates well with published results. As such, it was concluded that the use of MMSB sensing system embedded in OSIM iDesire massage chair is feasible in assessing the change in BVP.

With the success of this study, this development has also been implemented onto an embedded processor platform to automate the signal acquisition, filtering, signal recognition and statistical analysis. The developed product was found to produce measured results with 98% correlations to the laser Doppler measurements. Currently, the production prototype based on this development has been completed.

4 Relaxation assessment using heart rate variability

4.1 Introduction

Heart rate variability (HRV) is the change in the period between consecutive heartbeats over time. Such a change is predominantly dependent on the extrinsic regulation of the heart rate (HR). For healthy adult, the normal variability in HR is due to autonomic neural regulation of the heart and the circulatory system [11]. Due to external predictable and unpredictable stimuli, HRV is used as a measure of the ability of the heart to adapt to these changing circumstances. As such, analysis of HRV will allow the assessment on the overall cardiac health and the state of the autonomic nervous system (ANS), which is responsible for regulating cardiac activity. The ANS consist of the sympathetic nervous system (SNS) and parasympathetic nervous system (PNS) branches. The balancing action of the SNS and PNS branches of the ANS controls the HRV. Increased SNS or diminished PNS activity results in cardio-acceleration. Conversely, a low SNS activity or a high PNS activity causes cardio-deceleration. The degree of variability in the HR provides information about the functioning of the nervous control on the HR and the heart's ability to respond to stressors.

Stressors are often associated with an increase in sympathetic cardiac control, a decrease in parasympathetic control, or both. Associated with these reactions is a frequently reported increase in low frequency (LF, centred around 0.1 Hz, or 0.04–0.15 Hz) heart rate variability, a decrease in high frequency (HF, 0.12 or 0.15–0.4 Hz) power, and/or an increase in the LF/HF ratio. Aspects of this general response pattern have been reported for: (a) acute laboratory psychological or cognitive stressors such as mental arithmetic, reaction time tasks, Stroop interference task [12], or speech stress [13]-[17]; (b) real-life acute stressors such as college examinations [18], earthquakes [19], as well as typical day-to-day hassles [20]; and even (c) the level of chronic perceived stress associated with trait anxiety [21]. The study on the autonomic effects of acute laboratory stressors [13], for example, reported a stress-related decrease in HF heart rate variability (LF was not quantified), associated with a significant reduction in parasympathetic cardiac control and an increase in sympathetic control, as revealed by selective pharmacological blockades of the autonomic branches.

In this section, research and development on the use of HRV for relaxation assessment will be discussed together with implementation of an embedded system to quantify the relax level of a subject after going through a massage. In this experiment, the stimulus is the OSIM iDesire Massage chair and HRV is acquired using MMSB based sensing system. The directional change in HRV will be correlated to the SNS and PNS response to the stimulus and will be used as a surrogate marker to indicate the mental state (e.g. relax level) of the subject.

4.2 Experimental overview

Using the optimal sensor mounting on the heel and signal filtering outlined in section 3.2 of Chapter 6, this section of the research work will focus on the acquisition of HR data for HRV analysis on the heel. In addition, design and implementation of optimal sensor mounting for MMSB acquisition on a wrist structure / strip will be discussed. This work is part of the commercialization development based on the needs of the commercial company. As such, an overview of the sensors and magnets mounting will be discussed where limited details will only be presented in this section due to disclosure limitations.

4.2.1 System setup for human trial

In the system setup used for the trial, a wrist structure is designed as shown in Figure 119. An array of 3 sensors and 6 magnets are configured such that the magnets will produce a uniform magnetic field around the wrist. The sensors array will pick up the magnetic disturbance of the blood flowing through the uniform magnetic field as described by the principle of MMSB.

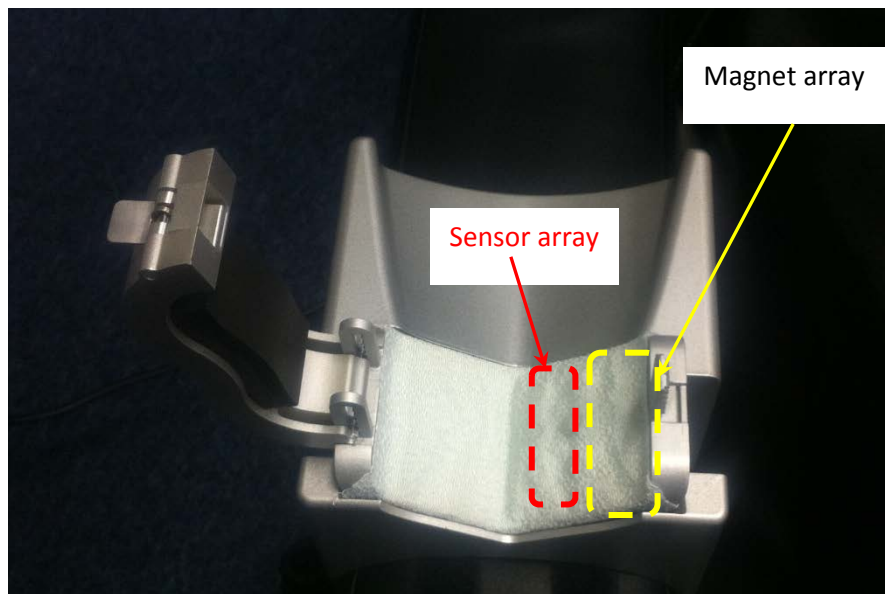


Figure 119 Sensors and magnets concealment on wrist structure

Using the sensors and magnets configuration as shown in Figure 119, once the wrist is aligned to the edge of the wrist structure, MMSB signals can be acquired for varying sizes of the wrist profile as shown in Figure 120. As such, MMSB signal acquisition is reliable and is independent on the user's familiarity of the system. The locking mechanism implemented in the wrist structure will help to keep the alignment of the sensors and magnets allowing MMSB signals to be acquired reliably for 2 minutes with minimum motion artefacts. The amount of time required for MMSB signal acquisition

is based on the recommendations for HRV measurements by the Task Force of The European Society of Cardiology and The North American Society of Pacing and Electrophysiology [26].

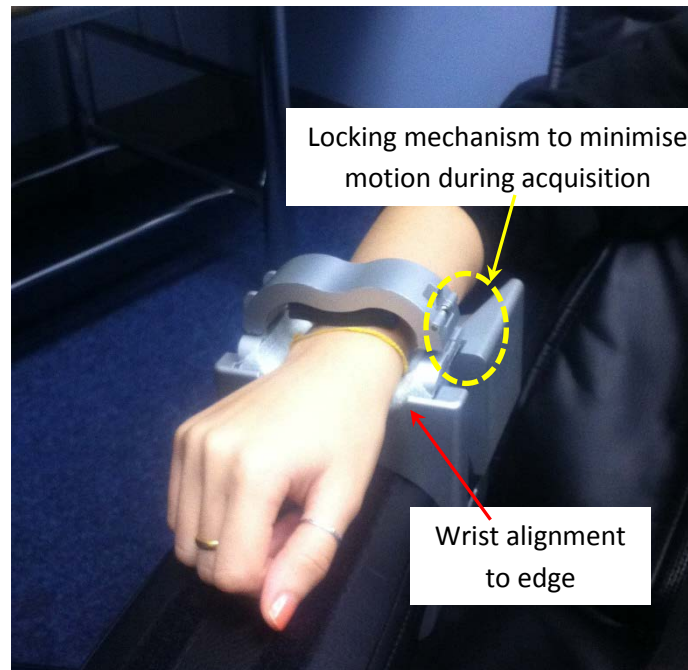


Figure 120 Placement of user wrist on wrist structure for MMSB acquisition

4.3 Setup and procedure used in trial

The subjects selected for this trial are healthy adults aged 19-55 years old. A total of 40 subjects, 20 females and 20 males, were randomly selected.

The control variables implemented for this trial are rest time for each subject before trial, ambience, stimulus (massage sequence) and time interval between each massage.

When the subject reports for the trial, each subject is given 5 minutes of rest time before the start of the experiment. This interval is selected as subjects arriving at the site of measurements maybe exposed to different level of physical (e.g. walking, taking the stairs, brisk walking, etc). For a typical healthy adult without going through excessive exercise, a period of 5 minutes is sufficient to achieve a stable (resting) physiological state for measurements to start.

During each of the measurements, the subject is exposed to similar ambience such as temperature (fixed at 23-25°C), light (fixed with use of yellow table lamp) and music (using a relaxation music provided by OSIM International Pte Ltd).

The stimulus for this experiment is the massage chair. However, in order to ensure all the subjects are exposed to a similar control variable, the massage sequence is set to be the same for all the

subjects with similar intensity. In addition, each measurement is not repeated on the same subject within a 24 hours interval. In this way, the measured variables will be independent of the each trial.

The measured parameters for this experiment are average Heart Rate (Pre and Post Massage), Relaxation Level (Pre and Post Massage using HRV) and the general state of health (e.g. fever, body aches, etc).

The experiment begins with the subject comfortably rested on the massage chair inclined at an angle of 30-40 degrees from the floor horizon. The temperature, light and music ambience are fixed as described above. Data acquisition is done using the wrist structure for a period of 2 minutes or 120 heart pulses. Using the acquired data, the pre-massage indication is computed together with the heart rate of the subject. Both measured variables are recorded for analysis in latter section.

Maintaining the same ambience, the stimulus of the experiment (i.e. massage) is applied on the subject for a period of 15 minutes. Upon the completion of the massage, data acquisition is completed using the wrist structure for a period of 2 minutes or 120 heart pulses. Using the acquired data, the post-massage indication is computed together with the heart rate of the subject.

4.4 HRV analysis using MMSB signals

In Chapter 2, section 7, instantaneous HR acquired using MMSB has been statistically validated to be accurate with reference to a gold instrumentation heart rate acquisition system, ECG. In this section, instantaneous HR will be extracted from MMSB signals for HRV analysis.

Based on the guidelines provided by the Task Force of The European Society of Cardiology and The North American Society of Pacing and Electrophysiology [26], HRV can be analysed using both the time-domain and frequency-domain approach.

Time domain HRV analysis are statistical measurements based on HR data collected over 24 hours (at least 20 minutes is recommended). In a continuous electrocardiographic (ECG) recording, each QRS complex is detected, and the so-called normal-to-normal (NN) intervals (that is all intervals between adjacent QRS complexes resulting from sinus node depolarisations), or the instantaneous heart rate is determined. Parameters that standard deviation of all NN intervals (SDNN) in seconds, square root of the mean of the sum of the squares of differences between adjacent NN interval (RMSSD) in milliseconds (ms), number of adjacent NN intervals differing more than 50 ms. (NN50 count), percentage of difference between adjacent NN intervals differing more than 50 ms. (pNN50 %), the integral of sample density distribution of RR intervals divided by the maximum of the

density distribution (RR triangular index) and baseline width of the minimum square difference triangular interpolation of the maximum of the sample density distribution of RR intervals in seconds (TINN). Standard deviation (STD) of the mean heart rate per minute is also computed.

In the frequency domain, HRV can be analysed by looking at the different frequency components of the frequency domain waveform. Three main spectral components are distinguished in a spectrum calculated from short term recordings of 2 to 5 minutes [27]-[31]: very low frequency (VLF), low frequency (LF), and high frequency (HF) components. The two main frequency components that represent ANS activity are the LF components (0.04 to 0.15Hz) and the HF components (0.15 to 0.4 Hz). Frequency domain measures confirm that the LF and HF oscillatory components are relative indices of cardiac sympathetic and vagal activity respectively and HF and RMSSD indicate parasympathetic activity [23]-[25].

In order to satisfy the consumer needs of stress assessment within a short period of time (i.e. 2 minutes or less), the frequency domain analysis of HRV using MMSB sensing is selected. The flow chart summarizing the steps for recording and processing of MMSB signal for frequency domain HRV analysis is illustrated in Figure 121.

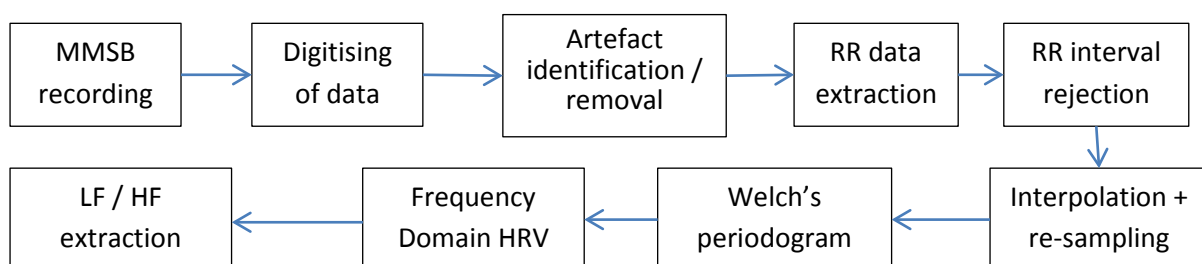


Figure 121 Steps for recording and processing MMSB signal for frequency domain HRV analysis

After the QRS complex occurrence times have been identified in the MMSB signal, the RR interval will be extracted as shown in Figure 122. The inter-beat intervals or RR intervals are obtained as differences between successive R-wave occurrence times. That is, the n 'th RR interval is obtained as the difference between the R-wave occurrence times $RR_n = t_n - t_{n-1}$. In some context, normal-to-normal (NN) may also be used when referring to these intervals indicating strictly intervals between successive QRS complexes resulting from SA-node depolarization [26]. In practice, the NN and RR interval provides the same values. Therefore, in this section, the RR interval is used for HRV derivation.

Upon artefact identification and removal, extraction of the RR interval is done as the first step for HRV signal pre-processing. This is followed by for RR data clean up (i.e. rejection) and interpolation

or re-sampling as shown in Figure 123. Using FFT on the RR interval, frequency domain HRV analysis is obtained. Each of these steps will be discussed in the following sections.

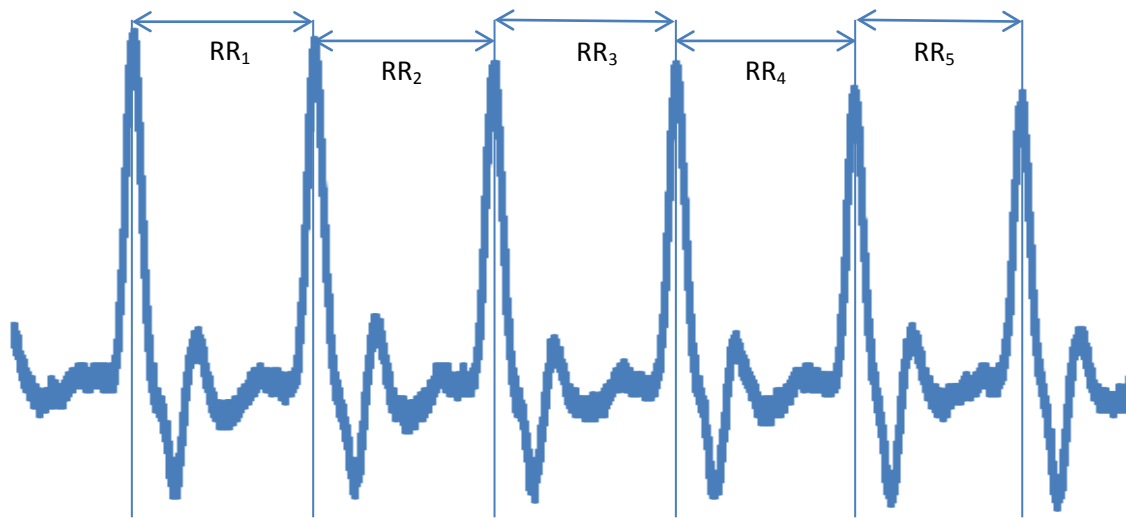
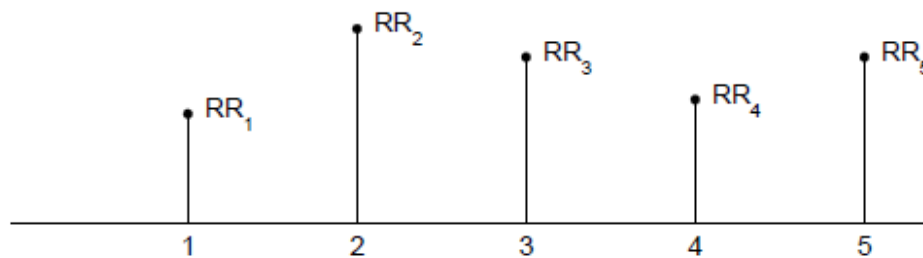


Figure 122 Derivation of RR interval from MMSB signals

RR interval tachogram



RR interval series (with two possible ways of interpolation)

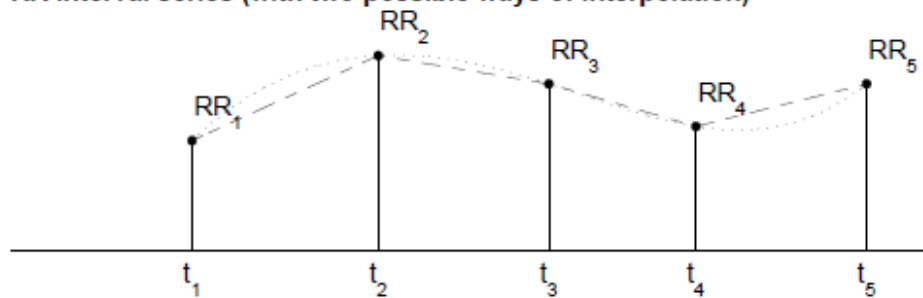


Figure 123 Plotting of RR interval tachogram and interpolation of the RR series for HRV derivation

4.4.1 Pre-processing of HRV time series

Artefact in the RR interval time series may interfere with the analysis of the HRV signals. The artefacts within HRV signals can be divided into technical and physiological artefacts. The technical artefacts can include missing or additional MMSB waveform detections and errors in peak-wave occurrence times. These artefacts may be due to measurement artefacts or errors in the

computational algorithm for MMSB recognition and peak extraction. The physiological artefacts, on the other hand, include ectopic beats and arrhythmic events. Such artefact is more difficult to remove and is mitigated through the selection of young and healthy adults for the trial.

In order to avoid the interference of such artefacts, the MMSB recording and the corresponding event series, sections of the MMSB waveform will be manually checked for artefacts to ensure that the analysis period are artefact-free [26]. In the event that the amount of artefact-free data is insufficient, the nonlinear predictive interpolation method is used to reduce these artefacts [33]-[35].

Another common feature that can alter the HRV analysis significantly is the slow linear or more complex trends within the analysed time series. Such slow non-stationaries are characteristic for HRV signals and should be understood before the analysis. The origins of such non-stationaries in HRV are discussed in [36]. The slow non-stationary trends from the HRV signal is removed using first order de-trending [37][38] model so as to provide smoothness prior to HRV PSD extraction. In this measurement, the moving average method is implemented to remove the slow non-stationary trend in HRV signal.

4.4.2 *FFT of HRV time series*

In the frequency-domain methods, a power spectrum density (PSD) estimate is calculated for the RR interval series. The regular PSD estimators implicitly assume equidistant sampling and, thus, the RR interval series is converted to equidistantly sampled series by interpolation methods prior to PSD estimation. In this section, a cubic spline interpolation method is used [32]. This is followed by the Fast Fourier Transform (FFT) using the Welch periodogram for PSD estimation.

The Welch periodogram is a method by which a large time-sampled waveform can be frequency-transformed by partitioning the data into shorter segments, transforming each segment, then, averaging the results over all the segments to create a composite frequency-space waveform.

The reasons for using the periodogram are as follows:

- the averaging of the individual PSDs allows for the reduction of variance errors
- it is logarithmically more efficient to take many shorter FFTs than one longer one which makes this technique attractive for processing long runs of data [39]

To properly create a PSD profile, values for three parameters must be judiciously selected:

- Window size or data segment

- Overlap size
- Number of frequency points

4.4.2.1 Window size/Data Segment

The data segment is a subset of the total number of data samples, N , as shown in Figure 124. The segment, in time samples, determines the resolution within frequency waveform. For coarse time spacing (i.e. large Δ), one will experience fine discrimination or resolution in frequency (i.e. $1/\Delta$) and vice-versa.

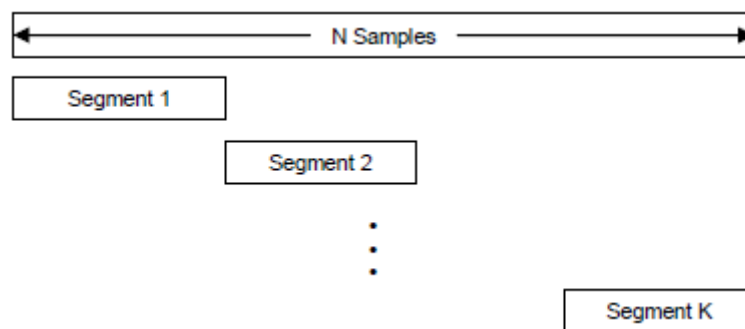


Figure 124 Segmentation of the N data sample source

In addition, data segments can introduce discontinuities at the segment boundaries or edges. These discontinuities manifest themselves as spurious frequency components in the transform space. To reduce these spurious components, the Hamming windowing function is applied in this experiment to taper the data down to zero at each end of the data segments.

4.4.2.2 Overlap size

The value assigned to the overlap parameter determines how many data segments can be obtained from the original sequence. For example, considering a 100 samples data sequence, if each data segment is 50 samples, then with no overlap, averaging can only occur between two PSDs. If overlap is set to 50%, then averaging over three PSDs is possible. More averaging reduces the error variance of the final PSD resolution. However, for averaging to reduce the error variance, it is assumed that the samples that get averaged are uncorrelated. If two segments are overlapped by a value approaching 90%, then the resulting PSD is nearly the same and averaging can no longer effectively reduce the error variance. In this experiment, the overlap size is set to 30%.

4.4.2.3 Number of frequency points (NFFT)

Increasing the number of frequency points affects the grid points on which the spectrum is evaluated. This parameter does not enhance physical resolution of the frequency spectrum but provides a finer grid on the frequency axis to observe the spectrum. Therefore, the higher the number of frequency points, the finer is the axis grid.

In choosing the NFFT parameter the following should be considered:

- For $NFFT \geq$ data segment, the FFT transform is zero-padded and all the samples of the data segment are applied to the transform
- $NFFT \leq$ data segment, some data samples are not applied to the FFT transform, which implies loss of frequency resolution

Therefore, in this experiment, the NFFT is set to 256 so that all the data samples collected over 2-3 minutes (i.e. 140-210 RR interval based on HR of 70bpm) are applied to the FFT transform without any loss of frequency resolution.

4.4.3 Power Spectral Density (PSD) analysis of HRV time series

The PSD plot obtained from the FFT by taking the absolute square sum of the complex FFT output. An example of the PSD plot is shown in Figure 125.

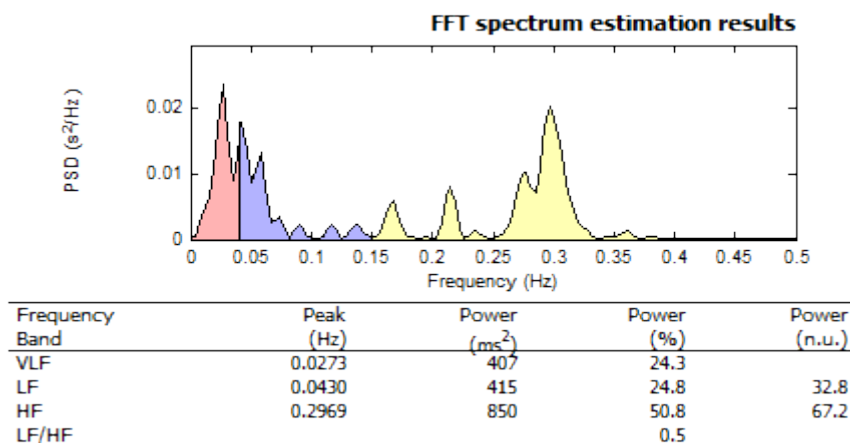


Figure 125 Example of a PSD plot for a typical RR data

The power for each frequency band (i.e. VLF, LF and HF) is calculated by taking the sum of all the points in the PSD plot for the specific band of frequency.

4.5 Qualification on effect of an external stimuli

Based on the association of relaxer with a decrease in low frequency (LF, centred around 0.1 Hz, or 0.04–0.15 Hz) power, an increase in high frequency (HF, 0.12 or 0.15–0.4 Hz) power, and/or a decrease in the LF/HF ratio, this section will outline the decision approach for the qualification on the effect of an external stimuli (i.e. massage) in relaxing the user.

In general, each measurement can be classified as parasympathetic or sympathetic dominance. The nominal range for healthy adults is extracted based on the age of the user extracted from [40]. For extreme conditions (i.e. the right top and left corner), data are classified as high energy and low energy state respectively. An exemplary illustration of this quantification is shown in Figure 126.

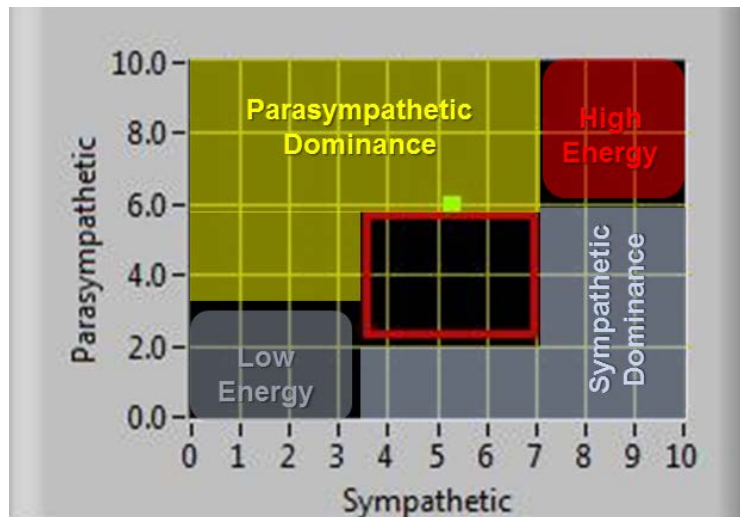


Figure 126 Illustration on directional change on HRV due to external stimuli

When an external stimulus is applied, the directional change of the HRV is an indication on the efficacy of the stimulus in relaxing or stressing the subject. Such a relative measurement can be quantified using the Equation 73.

$$\text{Equation 73} \quad \text{Directional Change, } D_{HRV} = \frac{\text{Post}\left(\frac{HF}{LF}\right)}{\text{Pre}\left(\frac{HF}{LF}\right)}$$

Similarly, the directional change can be illustrated as shown in Figure 127 where an increase in the sympathetic control is dominant with respect to a decrease in the parasympathetic control. Using data obtained from numerous conducted trials the classifications of stress, neutral and relaxation is shown in Table 27 (please note that the detailed data classifications is proprietary to the commercial company and the table is an exemplary illustration).

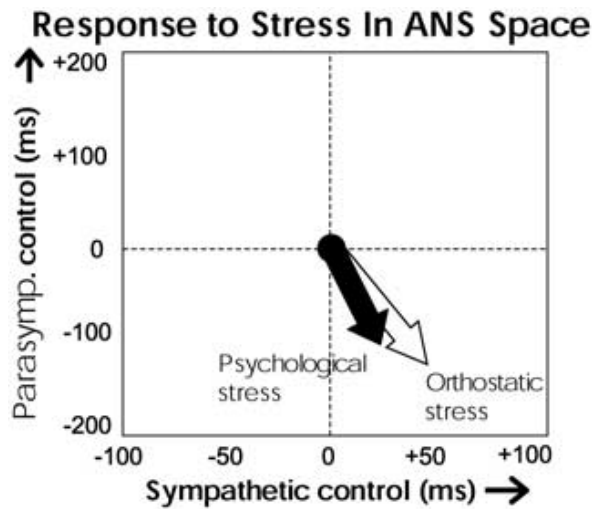


Figure 127 Illustration on directional change on HRV due to external stimuli

Level classifications	Indicative D_{HRV}
Stress	$D_{HRV} < 1.1$
Neutral	$0.9 \leq D_{HRV} \leq 1.1$
Relax	$0.9 < D_{HRV}$

Table 27 An exemplary classifications of DHRV as stress, neutral and relaxation

4.6 Statistical analysis of trial

The trial conducted for this section was to test for the concordance or inter-observer agreement (agreement of user perceived and BioS-HRV measured states). The statistical test used for the analysis of the results obtained is called Kappa correlation test [41] where Interpretation of the measured Kappa value can be summarized in Table 28. A value of 0 implies no agreement beyond chance, whereas a value of 1 corresponds to a perfect agreement between the two observers.

Concordance or inter-observer agreement	Kappa value
Poor agreement	= less than 0.20
Fair agreement	= 0.20 to 0.40
Moderate agreement	= 0.40 to 0.60
Good agreement	= 0.60 to 0.80
Very good agreement	= 0.80 to 1.00

Table 28 Interpretation of the measured Kappa value

There are situations where disagreements between observers may not all be equally important. For example, on an ordinal scale, a greater "penalty" will be applied if the two

categories chosen by the observers are farther apart. To account for these inequalities, Cohen (1968) introduced weights in the formulation of the agreement index leading to the weighted kappa coefficient. Although the weights are in general arbitrarily defined, those introduced by Cicchetti and Allison (1971) and by Fleiss and Cohen (1973) are the most commonly used. The former is linear and the latter has a quadratic form. Cohen (1968) showed that, under specific conditions, the weighted kappa coefficient is equivalent to the product-moment correlation coefficient. Moreover, Fleiss and Cohen (1973) and Schuster (2004) showed that the weighted Kappa with a quadratic weighting scheme is equivalent to the intra-class correlation coefficient.

In this analysis, weighted Kappa with a quadratic weighting scheme is used to analyse the agreement between the user perceived states of “Enjoyable,” “OK,” or “Painful,” and the relaxation states measured by the BioS-HRV system.

The user perceived states are obtained by asking the user on their experience after the stimulus had been applied. Users are allowed to express their perceived states without the knowledge of the relaxation state measured by the BioS-HRV system. As such, the correlation between the measured and perceived state are statistically independent.

To work out the Kappa statistics, Table 29, Table 30 and Table 31 outline the data transformation process where Equation 74 illustrates the Kappa statistics calculation.

		Observer 1			
		Relax	Neutral	Stressed	
Observer 2	Relax	a	b	c	Sum Row 1, SR1
	Neutral	d	e	f	Sum Row 2, SR2
	Stressed	g	h	i	Sum Row 3, SR3
		Sum column 1, SC1	Sum column 2, SC2	Sum column 3, SC3	Total, N

Table 29 Kappa statistics – Proportion agreement observed

		Observer 1		
		Relax	Neutral	Stressed
Observer 2	Relax	a/N	b/N	c/N
	Neutral	d/N	e/N	f/N
	Stressed	g/N	h/N	i/N

Table 30 Kappa statistics – Probability of proportion agreement observed (Po)

		Observer 1		
		Relax	Neutral	Stressed
Observer 2	Relax	(SC1/N)*(SR1/N)	(SC2/N)*(SR1/N)	(SC3/N)*(SR1/N)
	Neutral	(SC1/N)*(SR2/N)	(SC2/N)*(SR2/N)	(SC3/N)*(SR2/N)
	Stressed	(SC1/N)*(SR3/N)	(SC2/N)*(SR3/N)	(SC3/N)*(SR3/N)

Table 31 Kappa statistics – Probability of proportion agreement by chance (Pc)

Equation 74 Kappa statistics = $\frac{P_o - P_c}{1 - P_c}$

where $P_o = \sum \left(\frac{a}{N}, \frac{e}{N}, \frac{i}{N} \right)$, $P_c = \sum \left(\frac{SC1*SR1}{N^2}, \frac{SC2*SR2}{N^2}, \frac{SC3*SR3}{N^2} \right)$

The results obtained from the trial are tabulated as shown in Table 32, where the Po and Pc can be obtained using Table 33 and Table 34.

Subject Perceived Score	BioS-HRV Measured Score		
	Relax	Norm	Stress
Enjoyable	6	6	0
Ok	2	6	0
Painful	0	5	15

Table 32 Subject perceived score with BioS-HRV measured score

	Relax	Neutral	Stress
Relax	0.15	0.15	0
Ok	0.05	0.15	0
Painful	0	0.125	0.375

Table 33 Kappa statistics – Probability of proportion agreement observed (Po)

	Relax	Neutral	Stress
Relax	0.06	0.1275	0.1125
Ok	0.04	0.085	0.075
Painful	0.1	0.2125	0.1875

Table 34 Kappa statistics – Probability of proportion agreement by chance (Pc)

Using weighting based on seriousness of error (i.e. agreement expected by chance), the distance measurement is obtained in Table 35 with linear weightage as shown in in Table 36.

	Relax	Norm	Stress
Enjoyable	0	1	2
Ok	1	0	1
Painful	2	1	0

Table 35 Distance measurement based on Kappa statistics

	Relax	Norm	Stress
Enjoyable	1	0.75	0
Ok	0.75	1	0.75
Painful	0	0.75	1

Table 36 Weighted Kappa statistics (Quadratic)

Applying the product-moment correlation of Table 36 on the tabulated results for Po (Table 33) and Pc (Table 34), Table 37 and Table 38 are obtained respectively.

	Relax	Neutral	Stress
Relax	0.15	0.1125	0
Ok	0.0375	0.15	0
Painful	0	0.09375	0.375

Table 37 Kappa statistics – Weighted probability of proportion agreement observed (Po)

	Relax	Neutral	Stress
Relax	0.06	0.095625	0
Ok	0.03	0.085	0.05625
Painful	0	0.159375	0.1875

Table 38 Kappa statistics – Weighted probability of proportion agreement by chance (Pc)

Using the quadratic weightage for Kappa statistics, the observed Kappa value is calculated as 0.751. With reference to the qualification of Kappa value as shown in Table 29, the Kappa tests showed that there is “good” agreement between perceived state and stress state indicated by the BioS-HRV measurements. The “good” agreements indicate that the algorithm implemented in the Bio-HRV measurements is able to measure the perceived state of the user with sufficient accuracy as referred to the statistical tools used.

In conclusion, this section has demonstrated the ability of MMSB in providing HRV quantifications using statistical approach. The results on the statistical test to ascertain the concordance of user perceived state and MMSB acquired HRV state shows highly positive outcomes. Such positive demonstrates has demonstrated the accuracy of HRV in measuring the state of relaxation of the subject and is being developed as a commercial lifestyle product.

5 Conclusions on applications of MMSB

In this chapter, two applications were described and successfully developed using the MMSB as the method of measuring blood pulse information.

The remote heart rate monitoring application, titled Wireless Pulse Sense (WiPS), participated and won in the prestigious Tan Kah Kee Young Inventors Award 2007, Defence Science Category, in Singapore. WiPS was demonstrated to a panel of doctors and defence scientists. In addition, a short trial was conducted with the witness of the interview panel on 3 healthy subjects and the result was highly positive. With these success, WiPS was awarded the Silver Award [8], which is the highest award being given in this category of the competition for the past 8 years. The conferment of this award demonstrated the acceptance by the panel of doctors and defence scientists on the innovativeness and novelty of MMSB in acquiring blood pulse information. In addition, this development work was awarded S\$350,000 of funding to develop for deployment on a wireless ad-hoc network.

The application of MMSB in assessing the relaxation level of a subject using blood volume pulse and heart rate variability were implemented with the support of the needs of a commercial company. The success of this implementation has led to the development of the production model, which was completed at the point of this writing.

Through the applications developed in this chapter, the acceptance of MMSB, as a novel means to acquire blood pulse information, by the medical, defense and lifestyle industry were demonstrated. As MMSB describes a platform for pulse rate, blood flow and pulse transit time measurement, numerous lifestyle applications can be developed as MMSB is able to acquire signal through fabric, environmental contaminants and blood. Depending on the needs of the user of this technology, future applications such as energy expenditure and blood pressure measurements can also be developed for medical or research applications.

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Chapter 7: Conclusions, contributions and future work

1 Conclusions

The feasibility of achieving pulse rate[1]-[5], blood pressure[7] and blood flow[8] monitoring that is independent of the electrical or optical contact has been demonstrated in this dissertation. The MMSB phenomenon has been proven both practically[2] and mathematically[6][9] through physical measurements and simulations in a multi-physics environment respectively. In addition, physical measurements are also validated with comparison to other measurement devices such as pulse plethysmography, ECG[10][11], Laser Doppler Method and oscillometric blood pressure measurement[7].

Quantifications of MMSB measurements were successfully performed between gold standard medical instrumentation such as clinical Laser Doppler and ECG measurement instruments. For blood flow, PORH measurements were conducted on 6 healthy subjects using Laser Doppler and MMSB. Waveforms obtained from both measurement methods are highly correlated with differences of less than 4%. For heart rate (HR), 40 resting HR measurements were collected from 20 healthy subjects (aged 18-22) using concurrently ECG and MMSB acquisition with differences of < 1%. Statistical tests were used to study these measurements and it was concluded that this study supports the use of MMSB as a viable alternative to ECG for measuring resting HR for healthy individuals.

In addition, the mean arterial pressure (MAP) study was performed on 10 healthy subjects using the oscillometric based blood pressure lifestyle measurement device and the MMSB method. This study concluded the high correlation between the two methods of measurements with differences quantified to be < 8%.

In addition, two applications were developed to demonstrate the capability of MMSB in supporting low-cost lifestyle products, where ease of use and deployment can be incorporated into devices or integrated into furniture or beddings.

2 Contributions

This research has led to discovery of a novel sensing method of blood pulse and flow that is able to support measurements through fabric, environmental contaminants and body-fluids. Two major patents[1][2] and two divisional patents[3][4] were filed in the USA, Canada, Europe, India, China, Asia (Japan, Korea, Malaysia, Singapore) and Australia. All these patents were successfully defended in the Patent Cooperation Treaty (PCT) application with each receiving an International Preliminary Report on Patentability (IPRP) that has resulted in the grant of these patents in Singapore. In addition, the patent outlining the fundamental principle of MMSB[1] was granted by the USPTO in May 2012.

Together with these four patents are scientific publications made at six international conferences [5][6][7][8][9][10] and one international journal[11]. These publications are namely focused on the phenomena and science describing Modulated Magnetic Signature of Blood (MMSB); MMSB modelling in multi-physics software; measurements of blood flow using MMSB and; measurements of blood pressure using pulse transit time acquired using MMSB and ECG signals; validation of accuracy of MMSB in measuring heart rate with respect to a gold instrumentation using ECG.

Finally, two applications were successfully conceived and developed for continuous monitoring of pulse rate, HRV and blood flow. At the time of writing this dissertation, the production model for commercialization on the application of stress and relaxation measurements using blood flow and HRV had been successfully completed as a lifestyle product.

3 Future Work

There are two main areas of future work. First, the study of MMSB to acquire Mean Arterial Pressure (MAP) without the assistance of ECG signal will continue. This will enhance the existing work to support non-invasive; cuffless and occlusion free; electrical; and optical contact independent measurement of blood pressure possible. The potential of such a device will lead to the development of blood pressure monitoring devices that can be integrated into furniture and beddings for continuous monitoring with comfort and ease of usage. In addition, research and development on non-invasive measurement of pulse wave velocity (PWV) using MMSB is also in progress to assess the arterial stiffness and well-being of the subject.

Secondly, the application of the pulse rate obtained using the MMSB method for Energy Expenditure (EE) measurements. There are many researches being carried out to assess the EE of a subject. However most of the published work requires the measurement of VO_2 and VO_{2max} ^{*****}. Other published work applies heart rate measured from ECG or pulse oximetry for Energy expenditure measurement. This research work aims to develop a reliable EE measurement system using MMSB technology and physiological data from the subject.

In general, MMSB phenomenon offers promising possibilities of electrical or optical contact independent measurement of blood pulse, pressure and flow. This dissertation has presented the fundamental research and validation of the results, which is meant as the basis for broadening of potential application areas related to this technology.

***** VO_{2max} is the maximum amount of oxygen in millilitres, one can use in one minute per kilogram of body weight.

4 References

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Annex A - Datasheets

Annex A1 - Datasheet of INA118



INA118

Precision, Low Power INSTRUMENTATION AMPLIFIER

FEATURES

- **LOW OFFSET VOLTAGE:** 50µV max
- **LOW DRIFT:** 0.5µV/°C max
- **LOW INPUT BIAS CURRENT:** 5nA max
- **HIGH CMR:** 110dB min
- **INPUTS PROTECTED TO ±40V**
- **WIDE SUPPLY RANGE:** ±1.35 to ±18V
- **LOW QUIESCENT CURRENT:** 350µA
- **8-PIN PLASTIC DIP, SO-8**

APPLICATIONS

- BRIDGE AMPLIFIER
- THERMOCOUPLE AMPLIFIER
- RTD SENSOR AMPLIFIER
- MEDICAL INSTRUMENTATION
- DATA ACQUISITION

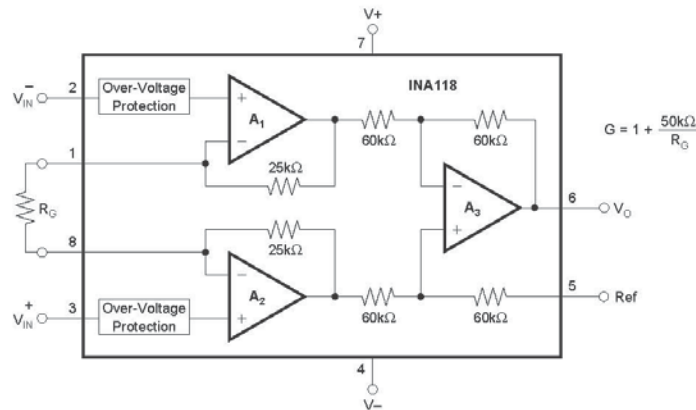
DESCRIPTION

The INA118 is a low power, general purpose instrumentation amplifier offering excellent accuracy. Its versatile 3-op amp design and small size make it ideal for a wide range of applications. Current-feedback input circuitry provides wide bandwidth even at high gain (70kHz at G = 100).

A single external resistor sets any gain from 1 to 10,000. Internal input protection can withstand up to ±40V without damage.

The INA118 is laser trimmed for very low offset voltage (50µV), drift (0.5µV/°C) and high common-mode rejection (110dB at G = 1000). It operates with power supplies as low as ±1.35V, and quiescent current is only 350µA—ideal for battery operated systems.

The INA118 is available in 8-pin plastic DIP, and SO-8 surface-mount packages, specified for the -40°C to +85°C temperature range.



International Airport Industrial Park • Mailing Address: PO Box 11400, Tucson, AZ 85734 • Street Address: 6730 S. Tucson Blvd., Tucson, AZ 85706 • Tel: (520) 746-1111 • Twx: 910-952-1111
Internet: <http://www.burr-brown.com/> • FAXLine: (800) 548-6133 (US/Canada Only) • Cable: BBRCORP • Telex: 066-6491 • FAX: (520) 889-1510 • Immediate Product Info: (800) 548-6132

APPLICATION INFORMATION

Figure 1 shows the basic connections required for operation of the INA118. Applications with noisy or high impedance power supplies may require decoupling capacitors close to the device pins as shown.

The output is referred to the output reference (Ref) terminal which is normally grounded. This must be a low-impedance connection to assure good common-mode rejection. A resistance of 12Ω in series with the Ref pin will cause a typical device to degrade to approximately 80dB CMR ($G = 1$).

SETTING THE GAIN

Gain of the INA118 is set by connecting a single external resistor, R_G , connected between pins 1 and 8:

$$G = 1 + \frac{50k\Omega}{R_G} \quad (1)$$

Commonly used gains and resistor values are shown in Figure 1.

The $50k\Omega$ term in Equation 1 comes from the sum of the two internal feedback resistors of A_1 and A_2 . These on-chip metal film resistors are laser trimmed to accurate absolute values. The accuracy and temperature coefficient of these resistors are included in the gain accuracy and drift specifications of the INA118.

The stability and temperature drift of the external gain setting resistor, R_G , also affects gain. R_G 's contribution to gain accuracy and drift can be directly inferred from the gain equation (1). Low resistor values required for high gain can make wiring resistance important. Sockets add to the wiring resistance which will contribute additional gain error (possibly an unstable gain error) in gains of approximately 100 or greater.

DYNAMIC PERFORMANCE

The typical performance curve "Gain vs Frequency" shows that, despite its low quiescent current, the INA118 achieves wide bandwidth, even at high gain. This is due to the current-feedback topology of the INA118. Settling time also remains excellent at high gain.

The INA118 exhibits approximately 3dB peaking at 500kHz in unity gain. This is a result of its current-feedback topology and is not an indication of instability. Unlike an op amp with poor phase margin, the rise in response is a predictable +6dB/octave due to a response zero. A simple pole at 300kHz or lower will produce a flat passband unity gain response.

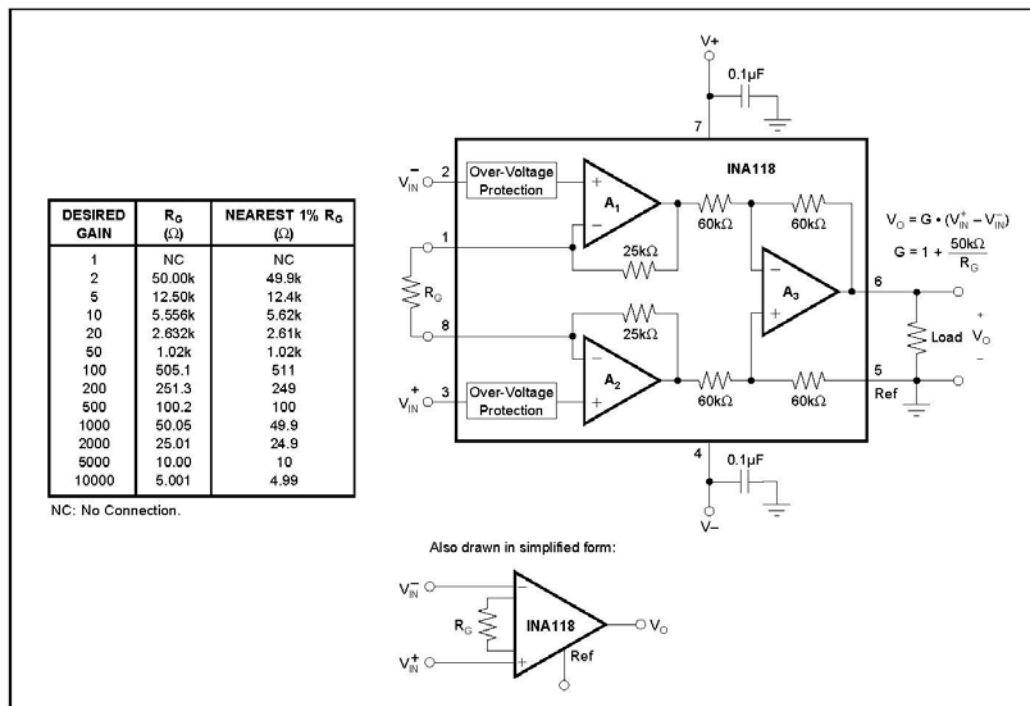


FIGURE 1. Basic Connections.

SINGLE SUPPLY OPERATION

The INA118 can be used on single power supplies of +2.7V to +36V. Figure 5 shows a basic single supply circuit. The output Ref terminal is connected to ground. Zero differential input voltage will demand an output voltage of 0V (ground). Actual output voltage swing is limited to approximately 35mV above ground, when the load is referred to ground as shown. The typical performance curve "Output Voltage vs Output Current" shows how the output voltage swing varies with output current.

With single supply operation, V_{IN}^+ and V_{IN}^- must both be 0.98V above ground for linear operation. You cannot, for instance, connect the inverting input to ground and measure a voltage connected to the non-inverting input.

To illustrate the issues affecting low voltage operation, consider the circuit in Figure 5. It shows the INA118, operating from a single 3V supply. A resistor in series with the low side of the bridge assures that the bridge output

voltage is within the common-mode range of the amplifier's inputs. Refer to the typical performance curve "Input Common-Mode Range vs Output Voltage" for 3V single supply operation.

INPUT PROTECTION

The inputs of the INA118 are individually protected for voltages up to $\pm 40V$. For example, a condition of $-40V$ on one input and $+40V$ on the other input will not cause damage. Internal circuitry on each input provides low series impedance under normal signal conditions. To provide equivalent protection, series input resistors would contribute excessive noise. If the input is overloaded, the protection circuitry limits the input current to a safe value of approximately 1.5 to 5mA. The typical performance curve "Input Bias Current vs Input Overload Voltage" shows this input current limit behavior. The inputs are protected even if the power supplies are disconnected or turned off.

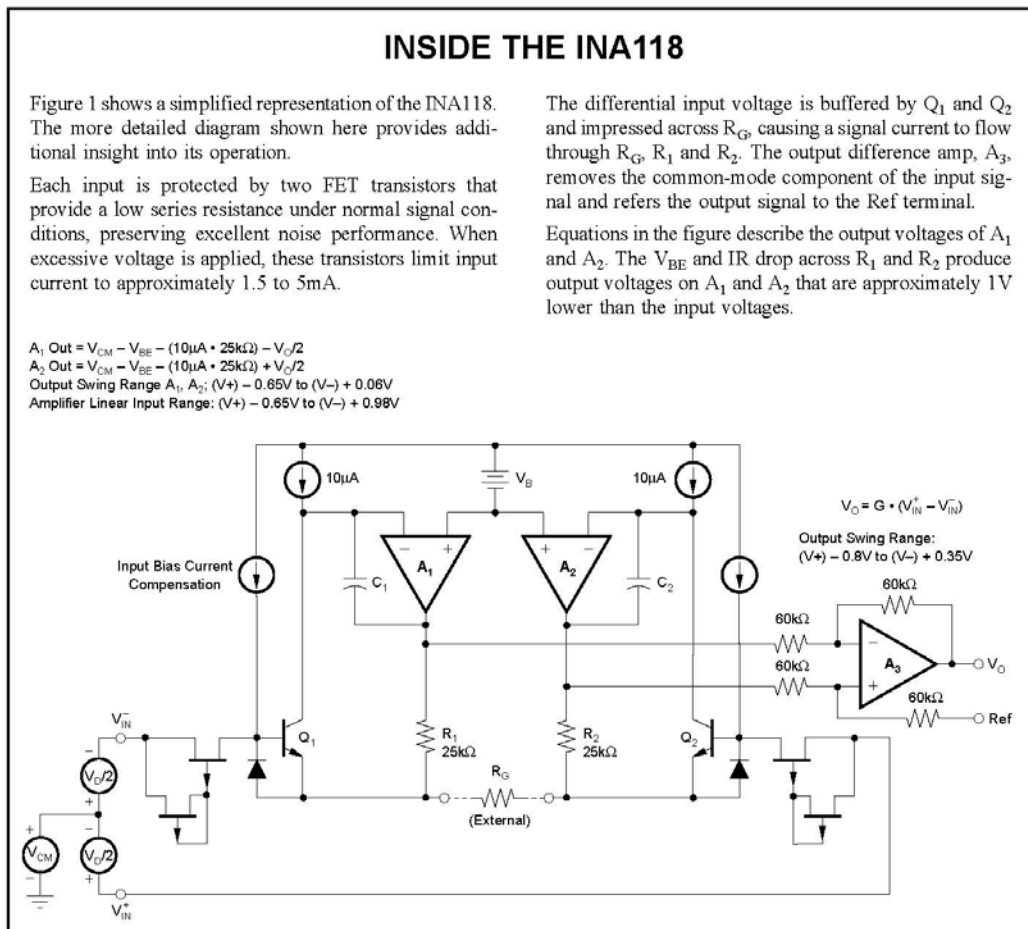


FIGURE 4. INA118 Simplified Circuit Diagram.

Annex A2 - Datasheet of INA321



**INA321
INA2321**

SBOS168D - DECEMBER 2000 - REVISED JANUARY 2006

microPower, Single-Supply, CMOS Instrumentation Amplifier

FEATURES

- **LOW QUIESCENT CURRENT:** 40µA/channel
Shut Down: < 1µA
- **HIGH GAIN ACCURACY:** G = 5, 0.02%,
2ppm/°C
- **GAIN SET WITH EXTERNAL RESISTORS**
- **LOW OFFSET VOLTAGE:** ±200µV
- **HIGH CMRR:** 94dB
- **LOW BIAS CURRENT:** 10pA
- **BANDWIDTH:** 500kHz, G = 5V/V
- **RAIL-TO-RAIL OUTPUT SWING:** (V+) - 0.02V
- **WIDE TEMPERATURE RANGE:**
-55°C to +125°C
- **SINGLE VERSION IN MSOP-8 PACKAGE AND
DUAL VERSION IN TSSOP-14 PACKAGE**

DESCRIPTION

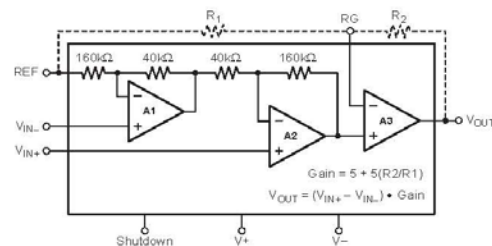
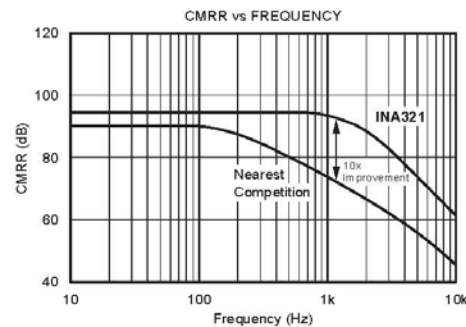
The INA321 family is a series of rail-to-rail output, micropower CMOS instrumentation amplifiers that offer wide-range, single-supply, as well as bipolar-supply operation. The INA321 family provides low-cost, low-noise amplification of differential signals with micropower current consumption of 40µA. When shutdown, the INA321 has a quiescent current of less than 1µA. Returning to normal operations within microseconds, the shutdown feature makes the INA321 optimal for low-power battery or multiplexing applications.

Configured internally for 5V/V gain, the INA321 offers exceptional flexibility with user-programmable external gain resistors. The INA321 reduces common-mode error over frequency and with CMRR remaining high up to 3kHz, line noise and line harmonics are rejected.

The low-power design does not compromise on bandwidth or slew rate, making the INA321 ideal for driving sample Analog-to-Digital (A/D) converters as well as general-purpose applications. With high precision, low cost, and small packaging, the INA321 outperforms discrete designs, while offering reliability and performance.

APPLICATIONS

- **INDUSTRIAL SENSOR AMPLIFIERS:**
Bridge, RTD, Thermistor, Position
- **PHYSIOLOGICAL AMPLIFIERS:**
ECG, EEG, EMG
- **A/D CONVERTER SIGNAL CONDITIONING**
- **DIFFERENTIAL LINE RECEIVERS WITH GAIN**
- **FIELD UTILITY METERS**
- **PCMCIA CARDS**
- **COMMUNICATION SYSTEMS**
- **TEST EQUIPMENT**
- **AUTOMOTIVE INSTRUMENTATION**



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APPLICATIONS INFORMATION

The INA321 is a modified version of the classic *two op amp* instrumentation amplifier, with an additional gain amplifier.

Figure 1 shows the basic connections for the operation of the INA321 and INA2321. The power supply should be capacitively decoupled with 0.1µF capacitors as close to the INA321 as possible for noisy or high-impedance applications.

The output is referred to the reference terminal, which must be at least 1.2V below the positive supply rail.

OPERATING VOLTAGE

The INA321 family is fully specified over a supply range of +2.7V to +5.5V, with key parameters assured over the temperature range of -55°C to +125°C. Parameters that vary significantly with operating conditions, such as load conditions or temperature, are shown in the Typical Characteristics.

The INA321 may be operated on a single supply. Figure 2 shows a bridge amplifier circuit operated from a single +5V supply. The bridge provides a small differential voltage riding on an input common-mode voltage.

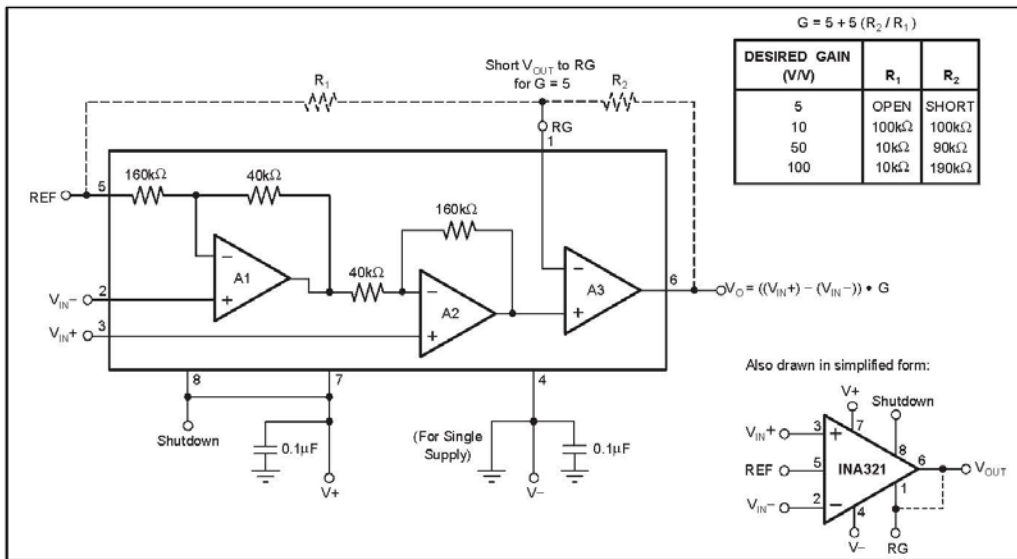


Figure 1. Basic Connections

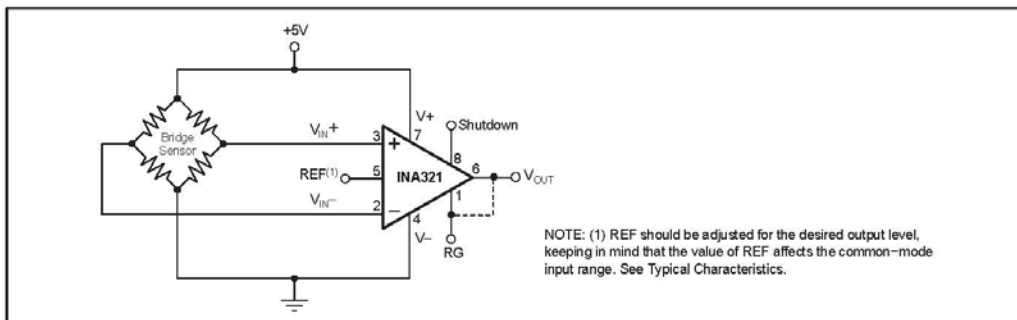


Figure 2. Bridge Amplifier of the INA321

SETTING THE GAIN

The ratio of R_2 to R_1 , or the impedance between pins 1, 5, and 6, determines the gain of the INA321. With an internally set gain of 5, the INA321 can be programmed for gains greater than 5 according to the following equation:

$$G = 5 + 5 (R_2/R_1)$$

The INA321 is designed to provide accurate gain, with gain error assured to be less than 0.1%. Setting gain with matching TC resistors will minimize gain drift. Errors from external resistors will add directly to the gain error, and may become dominant error sources.

INPUT COMMON-MODE RANGE

The upper limit of the common-mode input range is set by the common-mode input range of the second amplifier, A2, to 1.2V below positive supply. Under most conditions, the amplifier operates beyond this point with reduced performance. The lower limit of the input range is bounded by the output swing of amplifier A1, and is a function of the reference voltage according to the following equation:

$$V_{OA1} = 5/4 V_{CM} - 1/4 V_{REF}$$

(See Typical Characteristics for *Input Common-Mode Range vs Reference Voltage*).

REFERENCE

The reference terminal defines the zero output voltage level. In setting the reference voltage, the common-mode input of A3 should be considered according to the following equation:

$$V_{OA2} = V_{REF} + 5 (V_{IN+} - V_{IN-})$$

For optimal operation, V_{OA2} should be less than $V_{DD} - 1.2V$.

The reference pin requires a low-impedance connection. As little as 160Ω in series with the reference pin will degrade the CMRR to 80dB. The reference pin may be used to compensate for the offset voltage (see *Offset Trimming* section). The reference voltage level also influences the common-mode input range (see *Common-Mode Input Range* section).

INPUT BIAS CURRENT RETURN

With a high input impedance of $10^{13}\Omega$, the INA321 is ideal for use with high-impedance sources. The input bias current of less than 10pA makes the INA321 nearly independent of input impedance and ideal for low-power applications.

For proper operation, a path must be provided for input bias currents for both inputs. Without input bias current paths, the inputs will *float* to a potential that exceeds common-mode range and the input amplifier will saturate.

Figure 3 shows how bias current path can be provided in the cases of microphone applications, thermistor applications, ground returns, and dc-coupled resistive bridge applications.

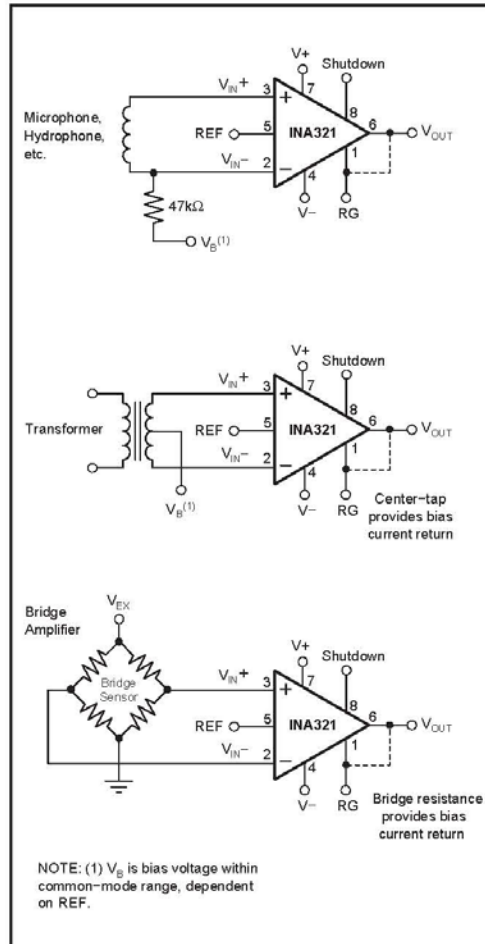


Figure 3. Providing an Input Common-Mode Path

When differential source impedance is low, the bias current return path can be connected to one input. With higher source impedance, two equal resistors will provide a balanced input. The advantages are lower input offset voltage due to bias current flowing through the source impedance and better high-frequency gain.

OUTPUT BUFFERING

The INA321 is optimized for a load impedance of 10kΩ or greater. For higher output current the INA321 can be buffered using the OPA340, as shown in Figure 4. The OPA340 can swing within 50mV of the supply rail, driving a 600Ω load. The OPA340 is available in the tiny MSOP-8 package.

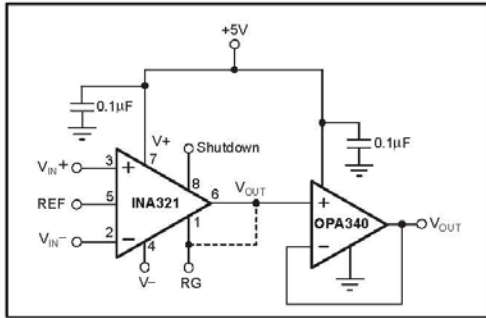


Figure 4. Output Buffering Circuit. Able to drive loads as low as 600Ω.

SHUTDOWN MODE

The shutdown pin of the INA321 is nominally connected to V+. When the pin is pulled below 0.8V on a 5V supply, the INA321 goes into sleep mode within nanoseconds. For actual shutdown threshold, see the Typical Characteristic curve, *Shutdown Voltage vs Supply Voltage*. Drawing less than 1µA of current, and returning from sleep mode in microseconds, the shutdown feature is useful for portable applications. Once in sleep-mode, the amplifier has high output impedance, making the INA321 suitable for multiplexing.

RAIL-TO-RAIL OUTPUT

A class AB output stage with common-source transistors is used to achieve rail-to-rail output for gains of 10 or greater. For resistive loads greater than 25kΩ, the output voltage can swing to within a few millivolts of the supply rail while maintaining low gain error. For heavier loads and over temperature, see the Typical Characteristic curve, *Output Voltage Swing vs Output Current*. The INA321's

low output impedance at high frequencies makes it suitable for directly driving Capacitive Digital-to-Analog (CDAC) input A/D converters, as shown in Figure 5.

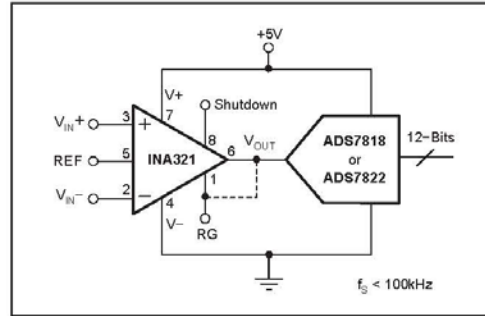


Figure 5. INA321 Directly Drives a Capacitive-Input, A/D Converter

OFFSET TRIMMING

The INA321 is laser-trimmed for low offset voltage. In the event that external offset adjustment is required, the offset can be adjusted by applying a correction voltage to the reference terminal. Figure 6 shows an optional circuit for trimming offset voltage. The voltage applied to the REF terminal is added to the output signal. The gain from REF to VOUT is +1. An op-amp buffer is used to provide low impedance at the REF terminal to preserve good common-mode rejection.

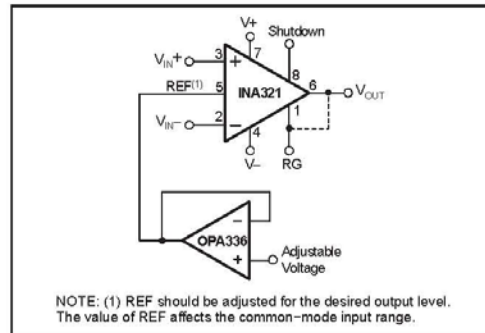


Figure 6. Optional Offset Trimming Voltage

INPUT PROTECTION

Device inputs are protected by ESD diodes that will conduct if the input voltages exceed the power supplies by more than 500mV. Momentary voltages greater than 500mV beyond the power supply can be tolerated if the current through the input pins is limited to 10mA. This is easily accomplished with input resistor R_{LIM} , as shown in Figure 7. Many input signals are inherently current-limited to less than 10mA; therefore, a limiting resistor is not required.

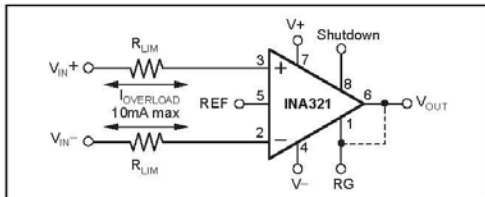


Figure 7. Input Protection

OFFSET VOLTAGE ERROR CALCULATION

The offset voltage (V_{OS}) of the INA321E is specified at a maximum of 500 μ V with a +5V power supply and the common-mode voltage at $V_S/2$. Additional specifications for power-supply rejection and common-mode rejection are provided to allow the user to easily calculate worst-case expected offset under the conditions of a given application.

Power-Supply Rejection Ratio (PSRR) is specified in μ V/V. For the INA321, worst-case PSRR is 200 μ V/V, which means for each volt of change in power supply, the offset may shift up to 200 μ V. Common-Mode Rejection Ratio (CMRR) is specified in dB, which can be converted to μ V/V using the following equation:

$$CMRR \text{ (in } \mu\text{V/V)} = 10[(CMRR \text{ in dB}) - 20] \cdot 10^6$$

For the INA321, the worst-case CMRR over the specified common-mode range is 90dB (at $G = 25$) or about 30 μ V/V. This means that for every volt of change in common-mode, the offset will shift less than 30 μ V.

These numbers can be used to calculate excursions from the specified offset voltage under different application conditions. For example, an application might configure the amplifier with a 3.3V supply with 1V common-mode. This configuration varies from the specified configuration, representing a 1.7V variation in power supply (5V in the offset specification versus 3.3V in the application) and a 0.65V variation in common-mode voltage from the specified $V_S/2$.

Calculation of the worst-case expected offset would be as follows:

$$\text{Adjusted } V_{OS} = \text{Maximum specified } V_{OS} + (\text{power-supply variation}) \cdot PSRR + (\text{common-mode variation}) \cdot CMRR$$

$$V_{OS} = 0.5\text{mV} + (1.7\text{V} \cdot 200\mu\text{V}) + (0.65\text{V} \cdot 30\mu\text{V}) = \pm 0.860\text{mV}$$

However, the typical value will be smaller, as seen in the Typical Characteristics.

FEEDBACK CAPACITOR IMPROVES RESPONSE

For optimum settling time and stability with high-impedance feedback networks, it may be necessary to add a feedback capacitor across the feedback resistor, R_F , as shown in Figure 8. This capacitor compensates for the zero created by the feedback network impedance and the INA321's RG-pin input capacitance (and any parasitic layout capacitance). The effect becomes more significant with higher impedance networks. Also, R_X and C_L can be added to reduce high-frequency noise.

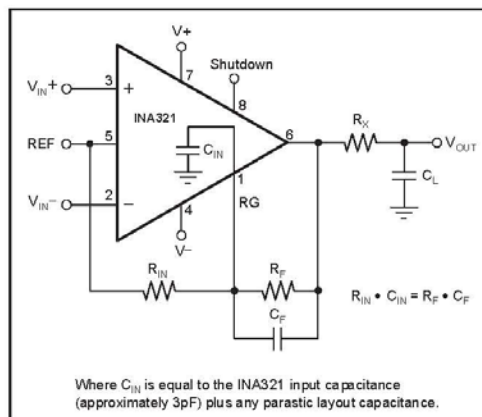


Figure 8. Feedback Capacitor Improves Dynamic Performance

It is suggested that a variable capacitor be used for the feedback capacitor since input capacitance may vary between instrumentation amplifiers, and layout capacitance is difficult to determine. For the circuit shown in Figure 8, the value of the variable feedback capacitor should be chosen by the following equation:

$$R_{IN} \cdot C_{IN} = R_F \cdot C_F$$

where C_{IN} is equal to the INA321's RG-pin input capacitance (typically 3pF) plus the layout capacitance. The capacitor can be varied until optimum performance is obtained.

Annex B – Comparison of methods to assess stress

Annex B-1: Table of Comparison on different methods of stress measurements

Title of Article	EEG*	Blood Flow	Heart Rate (HR) / HR Variability	Skin Temperature	Galvanic Skin Response	Blood Pressure	Pupil Diameter	Respiration Rate
1. Effect of Mental Stress on Coronary Flow Velocity Reserve in Healthy Men		X						
2. Measurement of Anxiety of Forearm Blood Flow		X						
3. Regulation of myocardial blood flow response to mental stress in healthy individuals		X						
4. Vasoconstriction and blood flow responses in visceral arteries to mental task in humans		X						
5. Flow dependence of forearm noradrenaline overflow, as assessed during mental stress and sodium nitroprusside infusion		X						
6. Head Level Blood perfusion under Acceleration		X						
7. Stress Detection in Computer Users Based on DSP of Noninvasive Physiological Variables		X		X	X		X	
8. Stress Recognition Using Non-invasive Technology		X		X	X		X	

Title of Article	EEG*	Blood Flow	Heart Rate (HR) / HR Variability	Skin Temperature	Galvanic Skin Response	Blood Pressure	Pupil Diameter	Respiration Rate
9. Detecting Exercise Induced Stress using Photoplethysmogram		X	X					
10. Level of Mental Stress May Help Doctors Predict Heart Attack		X	X					
11. Measuring Stress: A Guide for Health and Social Scientists by Sheldon Cohen, Ronald C. Kessler, Lynn Gordon, Lynn Underwood	X	X	X			X		
12. Effect of Mental Stress on Myocardial Blood Flow and Vasomotion in Patients with Coronary Artery Disease		X	X			X		
13. Measurement of Physiological Stress	X	X	X	X				
14. Development of a skin temperature measuring system for non-contact stress evaluation		X	X	X				
15. Analysis of skin-temperature variability compared to variability of blood pressure and heart rate			X	X				
16. Stress Response in Isolation			X		X			
17. Quantitative Assessment of Physiological Stress Effects Correlated with Mood States			X					X
18. VO2 Estimation Method Based on Heart Rate			X					X

Title of Article	EEG*	Blood Flow	Heart Rate (HR) / HR Variability	Skin Temperature	Galvanic Skin Response	Blood Pressure	Pupil Diameter	Respiration Rate
Measurement								
19. On Spectral Analysis of Heart Rate Variability during very slow yogic breathing			X					
20. The Analysis of Mental Stress using Time-frequency distribution of Heart Rate Variability Signal			X					
Percentage (%) of research using each technique to measure stress		33.33	30.77	12.82	7.69	5.13	5.13	5.13

* EEG is the accepted standard for stress measurement. Statistics shown indicates its usage as a reference for comparing with other methods for stress measurement

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RESUME

La mesure et le contrôle du pouls et du flux sanguin en continu sont d'importants paramètres pour l'évaluation de signes essentiels physiologiques sur la condition de santé d'un individu. Les dispositifs commerciaux existants, ainsi que les méthodes de recherche ou utilisées dans le milieu médical exigent un bon contact électrique ou optique pour obtenir cette mesure en continu. Pendant ces travaux de recherche, une méthode originale non invasive de mesure du rythme cardiaque fondée sur la perturbation localisée d'un champ magnétique au passage du flux sanguin a été développée, permettant l'acquisition des signaux à travers les vêtements, la transpiration, les salissures ou autres polluants dans l'environnement proche du capteur. Cette méthode est appelée la *Signature Sanguine par Modulation Magnétique* (MMSB) et les mesures ont été accomplies sur de multiples individus. Le système a été modélisé mathématiquement et simulé dans un environnement multiphysique, puis validé par l'utilisation des données expérimentales. Les résultats de mesure, en utilisant la méthode MMSB, pour le pouls et le flux sanguin ont été comparés et se trouvent bien corrélés, avec les résultats obtenus grâce à d'autres instruments. De plus, deux dispositifs ont été développés et sont en cours de commercialisation, pour des applications de vie quotidienne.

Mots Clés : pouls sanguin, pression sanguine, flux sanguin, bioMEMS, perturbations magnétiques

ABSTRACT

Continuous pulse rate, blood pressure and blood flow monitoring are important for the assessment of physiological vital signs as these are able to provide continuous feedback on the health condition of an individual. Existing commercial, medical and research methods to continuously acquire these physiological vital signs require good electrical or optical contact. During this research, a magnetic based sensing method, at room temperature, for blood pulse, flow and pressure is developed to achieve data acquisition through fabric, environmental contaminants and body-fluids. This method is named Modulated Magnetic Signature of Blood (MMSB) and physical measurements were conducted on multiple subjects, mathematically modelled and simulated in a multi-physics environment with verification through use of measurement data. Measurement results, using MMSB, for blood pressure and blood flow were compared, and found to be well correlated, with lifestyle device and medical research instruments respectively. In addition, two devices are developed, and are in the midst of commercialization, to support lifestyle applications.

Keywords : Blood pulse, pressure, blood flow, bioMEMS, magnetic disturbance
