A RESEARCH PROJECT PROPOSAL SUBMITTED IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE AWARD OF THE DEGREE OF MASTER OF SCIENCE IN APPLIED COMPUTING OF THE UNIVERSITY OF NAIROBI
DECLARATION

This thesis report is my own work and has not been presented for any degree award in any other university or institution of higher learning.

Signature _____________________  Date _____________________

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This thesis report has been submitted for examination with my approval as the university appointed supervisor.

Signature _____________________  Date _____________________

Prof Peter W. Wagacha
School of Computing and Informatics
University of Nairobi
DEDICATION

To our mothers. We salute you.

To all the women who are mothers and mothers to be.

We take our hats off to all those mothers who did not make it.

Our hearts go to their families and friends

I tip my hat to all the men and women doing research work to improve maternal care
ACKNOWLEDGEMENTS

With profound gratitude I acknowledge my supervisor Professor Peter W. Wagacha for the support and guidance he gave towards the completion of this project research thesis. I appreciate the patience, knowledge and encouragement he accorded to me through this journey far beyond his call of duty. I would also like to thank the University of Nairobi School of Informatics and Computing for their support and direction particularly the non-teaching staff and my fellow students.

I acknowledge the support and inputs I got from Kenyatta National Hospital and Mbagathi Hospital towards this research. I acknowledge the staff at the University of Nairobi, Science and Technology Park Maker Space for their feedback and inputs towards this research.

Finally, I acknowledge the support and encouragement I received for my family and friends.
ABSTRACT

Worldwide, maternal morbidity and mortality remains high with an estimated 830 women dying daily from preventable complications associate with pregnancy, with most of these cases occur in developing countries (WHO, 2017). The partograph has been recommend as a tool for use to help monitor labour progress to mitigate these risks. Nevertheless, completion of the partographs has proved to be evasive due to factors such as; staff shortage, inadequate training of staff and lack of supplies to monitor labour among others. These factors have thus resulted in a maternal mortality rate of 510 death for every 100,000 births in Kenya which is very high compared to the global ratio of 216 deaths for every 100,000.

In an effort to reduce maternal morbidity and mortality, this research seeks to study the current interventions used to mitigate maternal morbidity and mortality during the labour process and develop solutions around these findings.

Progress has been made in the area of maternal healthcare with an aim to reduce maternal mortality, related works that have been done before include; The partopen which is a digital pen used to fill a partograph form and is capable of giving time-based reminder with decision support. A second solution is the e-partogram. This is a monitoring and decision support electronic version of the partograph which focuses on connecting peripheral level providers to a central level supervisor. mlabour is another solution based on a mobile application which envelops an electronic partograph into workflows for recording patient details, updating and resolving maternal issues.

To develop the prototype, an awareness of the problem was studied with the use of secondary data. Tools used for measurement, under-staffing and overcrowding will be investigated with the help of questionnaires. We then built our prototype informed by the information collected. The prototype comprised of two main parts; the hardware which will be responsible for collection of the maternal vital signs, processing of the data and relaying it to the server which is the second part of the prototype. The server will be responsible for the storage of data received from the hardware as well as receiving and process request from a user.

The design of the prototype involved the selection of the resources to be used based on local availability. Once the resources were obtained the prototype was broken down to module and unit
tests carried out to test for correct functionality, after which the modules were integrated.

Partograph completion level was show to be less than 40%. Foetal heart rate and cervical dilation had the highest completion rate while urinalysis, moulding and amniotic fluid had the lowest.

The prototype would be useful in assisting nurses fill their partographs by recording vitals that a normally neglected or minimally recorded resulting in a better completion rate

Further work needs to be done to monitor foetal conditions as our study focused on maternal condition. Additionally, sharing of information among health facilities would further improve care.
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## DEFINITION OF TERMS

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>systolic blood pressure</td>
<td>Minimum blood pressure in between two heartbeats</td>
</tr>
<tr>
<td>Eclampsia</td>
<td>Convulsions occurring in a pregnant or puerperal woman, associated with preeclampsia, i.e., with hypertension, proteinuria, or edema</td>
</tr>
<tr>
<td>Effacement</td>
<td>The taking up or obliteration of the cervix in labour when it is so changed that only the thin external os remains.</td>
</tr>
<tr>
<td>Hypertension</td>
<td>A persistent blood pressure of 140 mm Hg systolic or 90 mm Hg diastolic.</td>
</tr>
<tr>
<td>Labour</td>
<td>The expulsion of conception through the vagina from the uterus</td>
</tr>
<tr>
<td>Meconium</td>
<td>The first tool passed out by the foetus that is made up of ingested materials consumed while in the uterus. These materials are the intestinal epithelial cells, amniotic fluid, lanugo, bile, water and mucus.</td>
</tr>
<tr>
<td>systolic blood pressure</td>
<td>Maximum blood pressure during one heartbeat</td>
</tr>
</tbody>
</table>
LIST OF ABBREVIATIONS

AIDS    Acquired Immunodeficiency Syndrome
EEPROM  Electronically Erasable Programmable Read Only Memory
HIV     Human Immunodeficiency Virus
HTML    Hypertext Markup Language
ICT     Information Communication Technology
IDE     Integrated Development Environment
IP      Internet Protocol
KNCHR   Kenya National Commission on Human Rights
PCB     Printed Circuit Board
RF      Radio Frequency
SDG     Sustainable Development Goal
SRAM    Static Random Access Memory
UI      User Interface
WHO     World Health Organisation
Wi-Fi   Wireless Fidelity
1 INTRODUCTION

1.1 BACKGROUND

1.1.1 KEY CHALLENGES IN MATERNAL HEALTHCARE

An estimated 830 women died daily from preventable difficulty associated with pregnancy in 2015 of which a majority occurred in developing countries (WHO, 2017). Statistics from the same report also shows that Kenya’s maternal mortality stood at 510 deaths for every 100,000 live births, well above the global maternal mortality ratio of 216 deaths for every 100,000 which in turn is well above the Sustainable Development Goals (SDGs) aim of 70 deaths for every 100,000 by 2030 (United Nations, 2017).

Approximately 99% of recorded maternal deaths occur in developing counties. Blood pressure that is high while pregnant (pre-eclampsia and eclampsia), bleeding excessively, infections (usually after childbirth), delivery difficulties and hazardous abortions are the major complication credited for approximately three quarters of maternal deaths. The remaining 25% result from or are associated with diseases like malaria and Human Immunodeficiency virus (HIV) / Acquired Immunodeficiency syndrome AIDS. Other factors adding to this statistic are the inability of some women to receive and seek care during labour due to; distance, lack of information, poverty, inadequate services and cultural practices (WHO, 2016).

In a study carried out across nine healthcare facilities in private, public and faith-based facilities ranging from a centre of health to tertiary hospital in four Kenyan provinces found that, low correct filling of the partograph which was blamed on the staff shortage within these facilities. In facilities where these tools were in use, records were incomplete with a 30% to 80% recording of contraction, 53% to 90% foetal heart rate recording and 70% to 97% cervical dilation recording. Minimal recording or documentation was done for the condition of the liquor, skull moulding and descent of the head as well as maternal parameters such as heart rate, blood pressure and urine composition. To worsen this situation, equipment for labour progress monitoring such as fetoscopes and blood pressure machines were in limited supply and some of those available were out of order (Qureshi et.al, 2010). This poor usage of the partograph was a result of shortage of staff, minimal/inadequate training and lacking of knowledge particularly with regards to interpreting findings, complexity of the form, negative attitudes, disputes among healthcare providers as to their
part in filling the form, and lack of role modelling by senior staff regarding the use, encouragement and application of the form (Qureshi et.al, 2010; Underwood, 2012). This poor use of the partograph is reflected in other hospitals in developing countries such as South Africa, Ethiopia and Nigeria with similar shortcoming recorded (Maphasha et al., 2017; Masika et al., 2015; Okokon et al., 2014; Ollerhead and Osrin, 2014; Yisma E et al. 2013).

A second study showed that in Kenya’s public health facilities providing delivery services only 36% had the bare minimum delivery infrastructure and equipment with remote and low-level facilities being poorly equipped. This infrastructure challenge in some hospitals is further complicated with the free maternal care offered by the government of Kenya thus resulting in; an increase in mothers well beyond the capacity of the maternity wards, mothers being required to vacate the facility earlier so as to create room for new arrivals as well as delivery on the hospital floors due to insufficient number of beds. Nurses are also overburdened, with almost all working extra-time and as much as 3 nurses assisting 20 to 49 mothers at one time (Bourbonnais, 2013; Underwood et al., 2012; Qureshi et.al, 2010).

With challenges as these being commonplace particularly in developing counties, the WHO endorses the used of partograph as an effective and affordable intervention tool for labour monitoring that if used properly could provide decision support to midwives by enabling them in detection emerging labour complications. This would prove beneficial more so in rural clinics with limited equipment, where the partograph can assist in early detection of complications giving health worked time to transfer the patient to a higher-level facility capable of handling these complications.

1.1.2 USE OF INFORMATION COMMUNICATION TECHNOLOGY (ICT) IN HEALTHCARE

There being an increase in use of information technology over the last decades and a half, more and more utilization of ICT in healthcare has been used to mitigate challenges facing the health system. Devices such as smart phones and tablets are starting to take the place of traditional monitoring and recording systems. Technology has also contributed to services being taken out of the health centres and into portable devices. Some of the impact of ICT on healthcare include increased accessibility, reduced cost of care, improved quality of care and increased education on health-related matters (Rudowski, 2008). However, various gaps can be seen in the maternal and child care sector as stated
above, these include a high mortality rate, limited infrastructures and human resources, poor service delivery, poor communication among health providers, limited knowledge on use of existing tools among others.

Some of the uses of ICT in health include; the electronic health record that is a central and productive tool in the retrieval of patient details as well as a health tool for data and population, mhealth which is a mobile healthcare device that enable healthcare providers and patients look at the healthcare processes on the go, telemedicine or telehealth is a tool that utilizes telecommunication technologies for remote diagnosis and treatment, Self-service kiosks for assisting hospital administrators check-in patients as well as payment for services, genome sequencing focuses on order of deoxyribonucleic acid (DNA) as well as diseases related to it, wireless communication with secure protocol for transmitting data within the hospital such as lab test, sensors and wearable technology such as heart rate monitor watches among others.

Some example of field use of these technologies includes home monitoring and telemedicine for the elderly. The system is applied in diseases such as cardiac failure, hypertension and diabetes and has been shown to assist in reducing the cost of healthcare by transmission of patient data over cellular networks (Rudowski, 2008). Apart from solving the poverty problem, the solution also looks into solving the problem of distance from a health centre. Transmission of ECG signals from the ambulance to invasive cardiology centre resulting in improved acute coronary syndrome diagnosis. (Rudowski, 2008). Teleradiology were images are transmitted to a location where radiologist on duty is available. Mobile phones and specialized tablets are also utilized as periodic reminder devices in busy facilities as well as for data entry and communication among healthcare providers.
encountered in the correct or complete use of the partograph as an intervention tool. These challenges include; shortage of staff and overburdened staff particularly in public facilities (which are further worsened with the introduction of free maternal healthcare provided by the Kenyan government), inadequate training and inability to interpret findings, complexity of the form, social factors and a shortage of labour monitoring equipment or no functional equipment.

Several technology-based solutions have been done in an effort to enhance the monitoring of labour progress with limited success. These solutions used time-based reminders that required the caregiver to attend to the patient of which in some cases was not possible due some of the challenges mentioned above such as staff shortages, inadequate training and form complexity.

This lack of proper monitoring of labour progress occasionally results in pregnancy related complication that may have been acted upon before the condition became critical or life threatening resulting in emergency actions which may include transfer of a mother to a higher-level hospital with the necessary equipment to handle the emergency.

Thus, this research study will seek to look at how we can leverage the use of microcontrollers and sensors technologies to achieve periodic data logging of maternal vital signs hence facilitating a higher completion rate of partographs by nurses enabling a better decision support system.

1.3 RESEARCH GAP

Several steps have been taken to decrease maternal morbidity and mortality, one mitigating solution is the utilization of the partograph a paper-based system for observing labour progress in which the caregiver periodically records the mother’s and foetus’ vitals. Interpreting of this tool has proved to be difficult in areas with caregivers who have limited knowledge of how to reading the partograph particularly in identifying emerging complications that may be developing.

To mitigate the challenge of limited skill level in using and interpreting the partograph, electronic solutions such as the partopen and e-partographs which serve as a training tool and an electronic based time reminder with decision support.

These tools have focused on solving various gaps found in maternal care which include monitoring of the vitals of both the mother and foetus and identifying developing problems. Providing decision
support to the caregivers and a training tool as well as using time-based reminders for periodic reading of these vitals.

Nevertheless, some gap in maternal care still exist. The gaps we are addressing are those of the time and human resource. Overcrowding in public facilities and understaffing makes it very challenging for caregivers to effectively monitor labour progress of all the mothers and within the specified periods that they need to be carried out in. Thus, this research seeks look at the possibility of automating the measuring and logging of maternal vital signs during the labour process and retrieval of these data to assist in the completion of a partograph as well as decision support. This would contribute to the efforts of mitigating maternal mortality during labour by leveraging information communication technology (ICT) in maternal care.

1.4 RESEARCH OBJECTIVES

The key objective of this research is to study how technology can be leverage to improve maternal healthcare.

The following are this research’s specific objectives:

- To review interventions that can decrease maternal mortality and morbidity
- To investigate existing solutions and how they are used.
- To develop a prototype of the data logger.
  - Integrate the various modules and microcontroller into one unit
  - To develop a software interface for data storage and retrieval
  - interface the data logger with the server
- Evaluate the system

1.5 RESEARCH QUESTIONS

- What activities are performed to ensure the best maternal care?
- How are maternal vital sign data monitored?
- How can ICT, microcontroller and sensor technologies be used to design a maternal vital sign monitoring?
- How does our solution compare to existing maternal vital sign monitoring solutions?
1.6 SCOPE
This research study will be limited to the following:

- Investigating and demonstrating whether and how microcontrollers and sensors can be used as a tool for collecting maternal vital signs
- Investigating existing tools used in hospitals in Kenya to measure maternal vital signs
- Designing and fabricating of a hardware prototype of the data logging solution.
  - The Data logger will focus on maternal vital signs alone
- Designing and implementing of the software that will capture and store the maternal vital signs.

1.7 JUSTIFICATION
This study seeks to enhance healthcare provided to expectant mothers by providing an auxiliary system for nurses and midwives by assisting them to record maternal vital signs. This service would result in a better completion rate of partographs forms thus better services in that complications can be detected early enough and appropriate action taken.

The process of completion of the partograph will be done through the use of information communication technology (ICT) to read data from the various sensors interpreted them to human readable information, storing them on a server and avail them to the health worker when requested. The successful use of ICT in this research could in turn encourage policies that favour the use of new and innovative technology in the field of maternal care opening the door to more innovative solution on ICT and automation that will improve efficiency in service delivery as well as cut cost of hiring extra staff and enabling the staff to perform other critical duties.
2 LITERATURE REVIEW

2.1 LABOUR AND ITS PROCESSES

2.1.1 INTRODUCTION

The Dorland's medical dictionary (2012) defines labour as the function of the female organism by which the product of conception is expelled from the uterus through the vagina to the outside world. The dictionary also divides the process of labour into four stages as explained:

- The first stage begins with regular contractions of the uterine and ends with the full dilation of the os uteri.
- The second stage two begins at the completion of stage one all the way to the expulsion of the infant.
- The third stage begins at the completion of stage two and ends at the expulsion of the placenta and membranes.
- Stage four is the period after delivery which lasts an hour or two, in which the uterine tone is established. The goal of this stage is to stop any profuse bleeding from the uterine atony (failure of the uterus to contract) and the cervical or vaginal lacerations.

2.1.2 MONITORED VITAL SIGNS

During labour, foetal assessments are performed to ensure that the foetal is well as well as detecting any signs that may indicate. The primary foetal assessments include the foetal heart rate (FHR) and patterns and character of the amniotic fluid. Irregularities in these assessments may be associated with abnormal foetal gas exchange and infection. FHR is assessed every hour during latent phase, half hourly during active phase and transition and quarter hourly during the second stage (Murray & McKinney, 2005).

Maternal assessments include monitoring of contractions, labour progress and vital signs. Labour progress is assessed to determine cervical dilation and effacement and foetal descent. Monitored maternal vital signs include blood pressure, heartbeat, temperature and respiration. Common values for these are; temperature 35.8°-37.3° C, heartbeat 60-100/min, respiration 12-20/min and blood pressure near baseline levels established during pregnancy. These signs are observed to identify the
start of hypertension and infection. The hypertension is in some cases a disorder specific to pregnancy or may be chronic, temperature of 38°C (100.4°F) and higher imply infection (Murray & McKinney, 2005). Temperature is taken after 4 hours and after 2 hours after membranes rupture or if elevated. While blood pressure, pulse and respiration are taken every hour (Murray & McKinney, 2005).

2.2 PARTOGRAPH

2.2.1 INTRODUCTION

Yisma E et al. (2013, pp 2) defines a partograph as a single page paper form that contains information about heart rate of the foetus, contraction of the uterine, drugs given and other important factors that could minimize descriptive notes. The partograph is a useful tool when used in an active labour ward with a large number of cases, but a small number of healthcare providers to screen for irregular labour. The partograph’s use removes the repeated recording of labour events. The tool enables the determination of deviations from normal labour progress, and assists in an intervention that is timely and proven.

The partograph is divided into 3 main parts:

- Foetal Condition: heart rate of the foetus, status of the amniotic fluid & moulding
- Progress of labour: cervical dilation, descent of the foetal head & uterine contractions
- Maternal condition: Heartbeat, blood pressure, temperature, urine, oxytocin & drugs given

The first WHO partograph covered the latent and active phases of labour. The latent phase covered a time period of 8 hours and the active phase began with a 3 cm dilatated of the cervix. The active phase sits between the alert line and an action line. These lines are drawn 4 hours apart on the partograph. This partograph’s used the principal that cervical dilation should be less than 1 cm/hour in the active labour (WHO, 1994). The modified WHO partograph was introduced to eliminate the disadvantages found in the first partograph. These modifications included few prolonged latent phases which was not usually associated with poor perinatal outcome and differentiating between the latent phase and false labour (Yisma E et al. 2013). The modified partograph does away with the latent phase and indicates 4 cm dilatation of the cervix instead of 3 cm as the beginning of the active phase.

Fistula Care (2011) lists benefits of the quality of service with proper partograph use that include enhance communication among providers, increase interaction between providers and the labouring
women, promote continuity of care across providers, and encourage teamwork.

2.2.2 FILLING THE PARTOGRAPH

Use of partograph begins in the active phase of labour. This is when the mother has regular contraction and the cervix is 4cm or more dilated. The main readings are the Cervix dilation, the descent of the head and hours in labour.

Patient information is filled at the top this are:
- The name of the patient
- gravida: The number of times the mother has been pregnant including those not carried to the end and the present pregnancy
- para: The number of above 24-week births (both viable and non-viable e.g. stillbirths). These are separated with a ‘+’ sign for example 2+0 would be 2 viable and no non-viable
- Hospital number
- Admission date
- Admission time
- Time of ruptured membrane or time elapsed since rupture of membrane

Foetal heart rate would be recorded every 30 minutes. Reading would usually be taken using a foetal doppler. To use this apparatus gel is usually placed on the mother’s stomach beforehand for ease of instrument movement and comfort and the doppler apparatus moved across the stomach as the healthcare giver locates the foetal heartbeat which would be displayed on the screen of the device. Another tool used in conjunction with a watch or clock is a foetal scope that is placed on the mother’s stomach and the foetus’s heartbeat timed.

Amniotic fluid colour is recorded at every virginal check-up and the table filled. ‘I’ indicates an intact membrane, ‘R’ membrane is raptured. If membrane is raptured, we use ‘C’ to indicate clear fluid, ‘M’ to indicate meconium stained and ‘B’ to indicate blood stained.

The extent to which the bones of the foetal skull are overlapping indicating the degree of compression the head is subjected to as it passes through the birth canal is referred to as moulding. Entry of moulding on the partograph is one using a scale from 0 to +1. ‘0’ indicates no moulding, ‘+1’ indicates that the bones are just touching each other, ‘+2’ indicated overlapping bones that are easily separated using finger pressure and ‘+3’ indicates overlapping bones that cannot be separated easily with finger
pressure.

Cervix dilation are recorded after 4 hours and marked with an ‘X’. The first plot is placed on the alert line. The action line is 4 hours away and parallel to the alert line. If labour progress is not following the expected course and the plot begins to approaches the action line, it would signal the need for action to be taken such as referrals before the action line is crossed. To record Cervix dilation, a vaginal examination is done by the caregiver.

Descent of the baby’s head is done at the same time as the vaginal examination and marked with an ‘O’ on the graph. This is assessed by abdominal palpation.

Hours in Labour and the time are recorded with the aid of a clock or watch.

Contractions are recorded every 30 minutes. This is done by counting the number of contractions within a 10-minute time period and record their length in seconds. For contractions count below 20 seconds dots are used to indicate this, for contractions between 20 and 40 seconds diagonal lines are used and if contractions last more than 40 seconds blacked out boxes are used.

If oxytocin is being used, the amount used is recorded as well as the volume of intravenous fluid it has been added to. The drops per minutes are recorded at the start and after every 30 minutes. Addition drugs given are recorded in their table

The mother’s pulse is recorded half hourly and marked with a dot. Her blood pressure is measured after 4 hours and marked with an arrow. The readings are usually measured with a digital blood pressure machine which gives readings of blood pressure and pulse rate on their display. Alternatively, a stethoscope and a mercury-based blood pressure machine are used.

The mother’s temperature is recorded every 2 hours. This reading is done using either a contact based or none contact-based thermometer.

Urine samples are collected and volume passed measured and test for protein and acetone content performed. Time urine was passed is also recorded.
### Figure 1: The modified WHO paragraph

<table>
<thead>
<tr>
<th>Name</th>
<th>Gravida</th>
<th>Para</th>
<th>Hospital number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date of admission</td>
<td>Time of admission</td>
<td>Ruptured membranes</td>
<td>hours</td>
</tr>
<tr>
<td>200</td>
<td>190</td>
<td>180</td>
<td>170</td>
</tr>
<tr>
<td>Fetal heart rate</td>
<td>140</td>
<td>130</td>
<td>120</td>
</tr>
<tr>
<td>Amniotic fluid Moulding</td>
<td>80</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cervix (cm) [Plot X]</td>
<td>7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Descent of head [Plot O]</td>
<td>6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hours</td>
<td>5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Contractions per 10 mins</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxytocin UIl drop/min</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drugs given and IV fluids</td>
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<td></td>
</tr>
<tr>
<td>Pulse</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>and BP</td>
<td>1</td>
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</tr>
<tr>
<td>Temp °C</td>
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</tr>
<tr>
<td>Urine</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>volume</td>
<td>acetone</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
2.3 RELATED WORKS

Several works of research and innovations have been done in the domain of maternal health utilising information communication technology (ICT) some of those related to this particulate work are:

2.3.1 PARTOPEN

To mitigate the challenge of skill level requirement found in using the partograph, the partopen was developed. The partopen is an electronic partograph solution usable as a training tool for medical students as well as an electronic time-based reminder with decision support. Its instruction manual are short audio recordings accessed by placing the pen on any of the text on the left side of the partograph. The pen is used for data entry on to a standard WHO partograph. The working principle of the pen is based on a Dot Position System which uses printed microdots arranged in specific patterns enabling the pen to establish its placement on a form (Underwood H et al., 2012). The electronic pen used to realise the Partopen solution includes a speaker for audio output, a microphone for audio input, a standard audio headphone jack, memory storage up to 8 gigabytes (equivalent to audio recording of 800 hours), a small display, a rechargeable battery, and a micro-USB connector for charging and data transfer.

2.3.2 EPARTOGRAM

The e-partogram (Saving Lives at Birth, 2011; jhpiego, 2009) is a handheld monitoring and decision support electronic version of the partograph designed for peripheral level providers. The device targeted the semi trained midwives. The e-partogram has feature contained in both the partograph and the partopen. The e-partogram was designed to change the perception of healthcare workers with respect to the partograph from being a burden on workflow to a helpful decision support tool.

The paper partograph was found to have various limitations that restricted its implementation. Some of the findings included the difficulty in learning how to fill the form correctly and lack of feedback thus failing to visualize the benefits to the user. The document was seen more as a recording device as opposed to a decision support tool, it is tedious to fill, difficult to interpret in real time and time consuming when obtaining all required measurements.

The challenges the e-partogram was looking to solve were:
• Provision of decision support
• Enable nurses to interpreter partographs easily
• Ease the process of data collection
• Better time management

Thus, the device enables efficient recording of labour, accurate determination of difficulties and a faster process of making choices. A sensory module is also included to partly automate the data collection process.

The design aim of the partogram was to make the device as easy as possible to use and interpret. Some of the design decision were hinged on the fact that the mobile phone is a ubiquitous device even in rural setting.

The e-partogram contains a telemedicine module that focus on connecting peripheral level providers to a central level supervisor enabling them to get support and guidance. The device's networking capacity is thanks to its communication module that utilises a sim card. The e-partogram also provides the ability of telemedicine between the peripheral level providers with the central level provider. With the devices networked the supervisor is able to monitored labour progresses with data transmitted to a web-server from the device and take action on problematic cases when they arise. The device also has an optional sensor belt that measure uterine contractions and can transmit text message data using a cell phone network.
2.3.3 MLABOUR

Schweers M et al. (2016, pp 2) define mlabour as a mobile application which envelops an electronic partograph into workflows for recording patient details, updating and resolving maternal issues. The primary work screen of the application is a list of patients that updates per minute and sorts on priority-bases which include: when the next examination is, risk level of patient, manual flagging by nurses for extra attention and if labour progress is progressing slow to the point of risk. Each patient’s major demographic and personal information is summed up on a tile, with images representing circumstances that are not usual: emergency during labour, history of high risk, examination overdue, or user set flags. The mobile application contains four major workflows which are:

- Patient registration, which collects identification data, obstetric history and high-risk factor screening.
• Record measurements, which are taken during an exam, this flow also includes information on the next check-up and an alarm is set.
• Notes and Flags: These are an alternative for information not recorded in the record measurement form which is more structured. A user is able to add or delete any tag showing a patient’s need of special attention. Details of the status are captured in a free text field.
• Resolution of Post-labour: this includes two forms, one for record delivery and another for recording transference to another doctor or facility

This digital solution of the partograph looked into improving on the standard partograph by increasing efficiency by generating graph instead of drawing by hand, error reduction by ensuring necessary measures are taken and validating measurements to expected range, increased usability by in-cooperating colour, animation and sound resulting in better data presentation and emphasis on abnormality.

2.4 FRAMEWORK
The works mentioned above were designed to meet various limitations encountered in the use of the partograph. The partopen uses a reminding system to alert nurses on pending reading to be taken as well as a training tool for medical students. The e-partogram contains feature found in both the partopen and partograph. Apart from having an alert system the e-partogram focuses on data sharing between peripheral level providers to central level supervisor. The mlabour is a simplified digital version of the partograph for use in mobile phone with features that assisting the midwife identify risks and slow labour progress.

These solutions require that the midwife periodically attend to a mother and record all necessary reading onto a physical or electronic partograph. Failure of which result in an incomplete partograph. A study by Underwood H et.al (2012; 2013) has shown that reminders and alert have proved to be a useful resource in the monitoring and management of mother during labour. The same study has also shown that due to limited workforce in the labour ward, midwives are not always able to attend to patients on a periodic basic (Qureshi et.al, 2010) has also shown that maternal vital signs are minimally recorded. This research seeks to file this gap caused by under-staffing and overcrowding by leveraging ICT, microcontrollers and sensors in implement a data logging solution for vital signs monitoring to assist nurses and midwives perform their duties through periodic measurements of blood pressure, respiration, heartbeat and temperature of mothers in labour.
The proposed solution will have two major parts; hardware and software. The hardware portion will constitute of various sensors for collecting vital signs, microcontroller for processing data from the sensors and a communication module for transmitting the data to a central server on which we will have a software for receiving the data from the hardware and compiling this into information that is useful to the nurses thus enabling them provide better service to mothers.

Figure 3: Conceptual Model
3 RESEARCH METHODOLOGY

3.1 INTRODUCTION

Design and creation research strategy was utilized as our research focuses on using technology in the development of a maternal vital sign data logger device.

Our study used an iterative process with five steps: knowing about the problem, proposing a solution, developing the solution, evaluation the solution and drawing conclusions (Vaishnavi & Kuechler, 2004).

Thus, the research study was organised as follows:

- Awareness of the problem
- Suggestion
- Development
- Evaluation
- Conclusion

3.2 AWARENESS OF THE PROBLEM

To get an understanding of the problem involved investigating the current condition within labour wards in health facilities including current interventions. This would involve interaction with nurses and midwives with the use of questionnaires, observation as well as looking at previously conducted researches. The objective here was to bring to light:

- Partograph completion rate
- Tools used for measurement
- How measurements are obtained
- Skills of caregiver
- Human resource - Overcrowding and under-staffing

Collection of data within any health facility in Kenya follows a strict protocol procedure. This
required the submission of our letter of approval from the school, a research protocol and questionnaire to the ethical board for consideration (See appendix D).

Once all requirements are met and approvals granted, we were required to brief all participants on the research and its’ objective as well as to ensure the confidentiality of the information collected (See appendices A, B and C)

Our Questionnaire was designed to collect data that gave us information on:

- Communication among the healthcare givers within a facility
- Communication among different facilities
- Skill level of the healthcare giver
- Availability of partographs
- Completion rate of the partograph
- Time management

A pilot of the questionnaire will be conducted to enhance reliability of the data collection tool. This will be conducted among 10 -15 randomly selected potential as well as gathering expert opinions. A final questionnaire will then be produced for this study.

Collection of measurement and tools used to take these measurements was done via observation as well as use of secondary data from other sources.

As mentioned above all correspondents were briefed on the research and what is expected of them as participants in the research. Our main target group for this research were those persons who work closely with the partograph, these include the midwives, nurses in labour wards and obstetricians.

3.3 SUGGESTIONS

Review of literature showed that 830 women died of complication related to pregnancy daily in 2015 and 216 died for every 100,000 live births globally (WHO, 2017), the WHO is in the forefront of promoting the use of the partograph as an effective and affordable intervention tool for labour progress checking. However, this has been difficult to implement particularly in developing countries with high ratio of maternal mortality.
Several challenges have played a role in the difficulty encountered in the correct or complete use of the partograph as an intervention tool in Kenya as well as other developing countries. These challenges include; shortage of staff and overburdened staff particularly in public facilities (which are further worsened with the introduction of free maternal services provided by the Kenyan government), inadequate training and inability to interpret findings, complexity of the form, social factors and a shortage of labour monitoring equipment or no functional equipment.

A number of technology-based solutions have been made in an effort to enhance the monitoring of labour progress with success in various aspects of the partograph. These solutions used time-based reminders that required the caregiver to attend to the patient of which in some cases was not possible due some of the challenges mentioned above such as staff shortages, inadequate training and form complexity.

Once these problems were identified through review of literature and data collection tools, a suggestion on how to solve the problem was proposed. A digital data logger to automatically monitor vital signs during labour process was seen to be the most viable option based on the information collected. Preparation of what needs to be done to achieve our goal was then developed. This was divided into 4 parts:

- Selecting the hardware components and relevant software
- Preparation of bill of materials
- Selection of software tools and supporting hardware
- Limitations of selections

The selection of hardware and software components was done on the basis of what was available locally. Ease of use of components and technical expertise also played a major part on what components were to be used. Once the components were identified information on the costing of each component was sought for after which a budget was created. The software interface on the computer side was then looked at, programming skill and operating system being used played a role on selection of some of the software used. As mentioned above the limitations encountered include; technical skill, availability of resource as well as time. Other limitation was on the basis of inability to record some of the condition due to lack of resource and others that require a visual inspection.
3.4 DEVELOPMENT

An ideal solution would to one that records all entries required in the partograph analyse progress of labour and automatically transfer the mother to a better health facility if complications are identified. However, this would require a great amount of technical expertise, resources and time that we do not have. Our proposed solution sought to implement some of the functions of an ideal solution. Hence our solution will focus on automation of some patient information, maternal vitals and foetal heart rate. This would be achieved by measurement of these entries and recording them into a digital storage where they may be retrieved on request by authorized personnel.

The design of the maternal vital sign data logging prototype was guided by the conceptual model as shown in Figure 3: Conceptual Model. The solution constitutes of two main modules which are the software module and the hardware module.

Components that make up the prototype will be selected based on functionality required, availability, ease of integration and expertise available. The next step is to develop the prototype device. This involved:

- Circuit design
- Assembly of prototype modules
- Setting up of a local web-server
- Development of the user interface
- Software hardware integration

The hardware portion of the prototype was further sub-divided into the various data collecting modules. Each of these modules require that a simple circuit be designed as well as writing it’s accompanying firmware program that would be loaded onto the chosen controller. These modules were:

- Blood pressure sensor
- Heart rate sensor
- Temperature sensor
- Fluid flow sensor
- Contraction sensor
- Foetal heart rate sensor
- Communication Module
- Computing module

Each of these modules would be individually tested after which they would be integrated into a single system for an integration test and ensure that everything is working as expected.

A web-server would be chosen based on the server’s operating system, these would include a web-server application, a database management system and a scripting language. The server will be used to store data received from the hardware module via its communication module as well as having a web-based user interface for users to communicate with the system. The user interface will enable the user to see records in a familiar readable format as well as input data into the database.

The final step of development will be the integration of the various hardware sub-modules, the software sub-modules and finally the hardware and software modules.

3.5 EVALUATION

To evaluate the prototype unit tests would be conducted on the various modules. For the hardware modules each sensory sub-module would be tested individually as well as the communication sub-module. This involved making sure that the sub-modules would functioning as expected and able to give correct readings as well as communicate with a Wi-Fi router. Once proper working of the sub-modules has been confirmed, the sub-modules would then be integrated into a final hardware prototype and tested as a single unit for proper functionality.

For the software portion of the prototype. A web interface would be created and customised so us to resemble a partograph form. It would then be tested on the various devices to check for correctness of display. For the second sub-module a script would be used to store and retrieve data from the database. Once proper working has been confirmed, it would be integrated to the web-server and data retrieval and display to check for proper working. This included checking that data from the hardware module were received correctly, user inputs were stored and processed correctly as well as correct display of user requests.
The prototype would then be presented to random participants for their views on the device. These participants include technical persons, those in the medical field and random persons.

Findings from other related research work would be analysed and patterns deduced and compared to our prototype.

3.6 ETHICAL CONSIDERATIONS

To conduct this research legal frameworks governing research activities typically require clearance from the relevant authorization bodies such as Kenyatta National Hospital – University of Nairobi Ethics and research committee (KNH-UoN ERC).
4 PROTOTYPE DESIGN AND ANALYSIS

The WHO endorses the use of partograph as an effective and affordable intervention tool for labour management that if used properly could provide decision support to midwives by enabling them in detection emerging labour complications. This would prove beneficial more so in rural clinics with limited equipment, where the partograph can assist in early detection of complications giving health worked time to transfer the patient to a higher-level facility capable of handling these complications.

The partograph is a paper-based system of monitoring labour progress where the health worker manually records entries into the form for each individual patient they are attending to. The nurse is then responsible for determining whether the labour is proceeding normally or complications may arise and take any necessary action.

4.1 PARTOGRAPH USE IN HEALTH FACILITIES

In a study in a sub county hospital in Kenya, where 177 partographs were analysed only 25.4% were complete, 63.3% had their admission details complete, 84.2% had the mother’s name complete, Cervical dilatation and descent were documented 74% and 54% respectively while the uterine contraction was at 49.7% and blood pressure at 46.9%. Factors that hindered completeness were workload, inefficient supervision, poor attitude, lack of motivation and supervision (Sigei, 2018).

In a second study across nine healthcare facilities partograph records were incomplete with a 30% to 80% recording of contraction, 53% to 90% foetal heart rate recording and 70% to 97% cervical dilation recording. Minimal recording or documentation was done liquor state, skull moulding and head descent as well as maternal parameters such as heartbeat, blood pressure and urinalysis. This poor usage of the partograph was attributed to shortage of staff, minimal/inadequate training and lacking of knowledge particularly with regards to interpreting findings, complexity of the form, negative mind-set, conflict among providers pertaining to the filling of the form, and lack of role modelling by senior staff with regards to utilization, promotion and application of the form (Qureshi et.al, 2010).

In a third study to see the effects of free delivery services on partograph utilization at Naivasha district hospital 88 partograph for the period July 2012 to December 2013 were assessed. Foetal heart rate was filled between 72.7% and 83%, 70% to 85% has adequate information on monitoring of cervical
dilation, descent and uterine contractions, blood pressure was recorded between 64.8% to 71.6%. Pulse was between 62.5% and 63.6%. Overall the completeness of partographs was less than 40% (Maina, 2016)

Kenyatta national hospital was selected as the field of research. The target demography are nurses in the labour ward and obstetricians. A questionnaire was developed as a data collection tools that will be used for gathering additional data such as nurse patient ratio, measuring tools as well as patterns and themes.

### 4.2 PROTOTYPE DESCRIPTION

An ideal solution would to one that records all entries required in the partograph analyse progress of labour and automatically transfer the mother to a better health facility if complications are identified. Our proposed solution sought to implement some of the functions of an ideal solution. They include monitoring of some of the maternal conditions by measuring them and recording them into digital storage where they can be retrieved when needed.

The design of the maternal vital sign data logging prototype was guided by the conceptual model as shown in Figure 3. The solution constitutes of two main modules which are the software module and the hardware module.

The prototype has two main modules as mentioned above. The main functions of the prototype are:

- To measure vitals of mother during labour
- Transmit the data to a web-server
- Store the measured data into a database
- Process queries on patient by health workers

The block diagram in figure 4 below visually describes these functions
4.2.1 TECHNOLOGIES

This section describes the hardware and software technologies used for the implementation of our prototype and their rational. The choice of hardware components was restricted to items that were available locally as well as affordable.

4.2.1.1 ARDUINO

The Arduino is an open source electronic hardware platform build around the 8bit and 32bit Atmel microcontrollers. It is a platform used by both novices and experts can develop and test prototype electronic project in a rapid manner. The platform comes with an integrated development environment (IDE) developed in processing and an Arduino programming language based on wiring. The platform also comes with a wide variety of libraries and supported hardware making prototyping simple.

The choice to use this platform was based on that we were familiar with the platform, its simplicity,
the low system requirements of the modules it used and that it had the necessary capability to perform all the function required to develop the hardware potion of the prototype

4.2.1.2 SENSORS

A variety of sensors were looked at with regards to the vital signs selected to be monitored, these are presented in table 1

<table>
<thead>
<tr>
<th>Vital</th>
<th>Sensor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood Pressure</td>
<td>U80EH automatic blood pressure monitor</td>
</tr>
<tr>
<td>Heart rate</td>
<td>U80EH automatic blood pressure monitor</td>
</tr>
<tr>
<td>Temperature</td>
<td>DS18B20 temperature sensor</td>
</tr>
<tr>
<td>Urine Volume</td>
<td>Flow meter</td>
</tr>
<tr>
<td>Contractions</td>
<td>Push button</td>
</tr>
</tbody>
</table>

Table 1: vitals and their sensors

The U80EH automatic blood pressure monitor was selected for measuring blood pressure and heart rate due to its availability in the local market as well as the ease of retrieving data from the device. The device also reduced our work load by interpretation of the blood pressure and heartbeat signals which was done by the device and their results simply sent to our controller for transmission.

4.2.1.3 COMMUNICATION MODULE

The communication module selected was the esp8266 module. This is a radio frequency (RF) module that uses Wi-Fi protocol to communicate with other devices. The module was selected as the communication device of choice due to its compatibility with the selected controller as well as its low cost.
4.2.1.4 MYSQL DATABASE

MYSQL database was selected for the database management system for managing patient data and their retrieval. It was selected due to its wide support and availability.

4.2.1.5 HTML AND PHP

Hypertext markup language (HTML) was used as the user interface (UI) and PHP was used to save patient information as well as retrieve and display results from the database.

4.3 PROTOTYPE DESIGN

As mentioned previously, the prototype constitutes of two main parts; the hardware and the software modules that constitutes our prototype solution.

4.3.1 HARDWARE

This sub-section describes the components of the hardware module and their circuit diagrams. Software used to achieve this were gimp which is an image manipulating software and eagle which is a schematic and printed circuit board design software. The firmware program for interpreting and transmitting data was written in the Arduino programming language as well as Atmel C++. The program was then loaded onto the microcontroller through the Arduino integrated development environment.

4.3.1.1 PROCESSING UNIT (ARDUINO)

An Arduino mega board was selected for use in the development of the prototype. The Arduino mega is a low power board equip with at Atmel atmega2560 8bit microcontroller that formed the brain of our hardware. This microcontroller has 256 kilobytes of flash memory, 4,096 bytes of electronically erasable programmable read only memory (EEPROM) and 8,192 bytes of static random access memory (SRAM) which is more than enough to store and execute our firmware. The Arduino mega also gave us access to 70 input and output pins that support digital and analogue signals as well as communication protocols enabling us connect all our sensors and communication module with plenty of pins to spare. The Arduino mega additionally has six timers enabling us to perform periodic routines critical to our prototype.
The microcontroller contains the firmware necessary for reading and interpretation of the voltages from the sensors of the various modules. This data was in-turn transmitted and stored in the server. The transmission was achieved with the use of a Wi-Fi radio frequency module. The microcontroller also acts as a time keeper with the help of a 16-megahertz quartz crystal thus facilitates for the periodic reading and transmission of measurements to the server.

4.3.1.2 BLOOD PRESSURE AND HEART RATE MEASUREMENTS

The measurements of mother’s blood pressure and heartbeat was achieved using a U80EH automatic blood pressure monitor. The U80EH automatic blood pressure monitor is a commercially available blood pressure monitoring system for home and clinic use. The monitor is able to measure the systolic blood pressure (maximum blood pressure within one heartbeat), diastolic blood pressure (minimum blood pressure in-between two heartbeat) and heart rate of a patient.
The U80EH automatic blood pressure monitor contains its own controller and algorithm for interpreting signals from its sensors. These are then stored into its internal memory as well as outputted to its debugging pins.

To get access to these pins the blood pressure monitor was disassembled exposing the printed circuit board. We then soldered jumper wires from the transmission (TXD) pin of the blood pressure monitor to the receive pin (RX1) of the microcontroller as well as shorting their ground terminals together. A third jumper wire was also connected between the to the start stop button of the blood pressure monitor and digital pin 12 of the Arduino mega so us to control the measuring and reading of blood pressure based on set predetermined values within the microcontroller’s firmware.
Figure 7: U80EH automatic blood pressure monitor debug pins

Figure 8: blood pressure monitor connected to an Arduino mega
4.3.1.3 TEMPERATURE MEASUREMENT

Temperature reading was achieved with the use of a DS18B20 digital temperature sensor. The sensor operation range is -55°C to 125°C with an accuracy of ±0.5°C and operation voltage range of 3 to 5.5 volts.

The temperature module was connected as shown in the figure below. Digital pin 10 from the Arduino was used to receive readings from the sensor. A 4700 ohms resistor between the supply voltage pin and signal pin was used as a pull-up resistor to enable accurate temperature conversion. The program for interpreting signals from the sensor into temperature readings was then loaded into the microcontroller.

![Figure 9: Temperature module circuit](image)

4.3.1.4 URINE VOLUME MEASUREMENT

The volume of urine was measured using a flow meter sensor. The flow rate sensor utilizes a hall effect sensor and a small magnet to measure flow. The magnet is placed on top of a small turbine wheel along the flow path of the fluid. Thus, when there is a fluid flow, the turbine rotates and magnet flux interference is registered by the hall sensor.

The flow sensor module was connected as shown in the figure below. Digital pin 4 from the Arduino was used to receive readings from the sensor. The program for interpreting the pulse signals from the sensor into volume was then loaded into the microcontroller.
4.3.1.5 CONTRACTIONS

Monitoring of contractions was implemented using a push button where a recording of the presses of the push button presses were captures. The microcontroller would then calculate the number of contractions per 10 minutes based on these button presses. The circuit was connected as shown in the diagram below with a pull-up resistor ensuring that the input received on digital pin 2 was stable. The accompanying program was then loaded onto the microcontroller.
Communication between the data logger and the server was done via the Wi-Fi protocol using the ESP8266 Wi-Fi module. The ESP8266 module consists of 8 pins used for various purposes as shown in the diagram below.

**4.3.1.6 COMMUNICATION MODULE**

Communication between the data logger and the server was done via the Wi-Fi protocol using the ESP8266 Wi-Fi module. The ESP8266 module consists of 8 pins used for various purposes as shown in the diagram below.

*Figure 11: Contractions simulation circuit*
The Wi-Fi module was connected as shown in the figure below the two diodes connected between digital pin 16 and the receiver of the Wi-Fi module are used to drop down the voltage as the Wi-Fi module works on a lower voltage than the Arduino. Digital pins 16 and 17 were used for serial communication between the module and Arduino. The Arduino was then loaded with a program to connect to the Wi-Fi module after which the module would then connect to a Wi-Fi network and data sent to the server.

*Figure 12: WIFI module pinout*
4.3.2 SOFTWARE

The software used for the prototype was setup on a Linux personal computer with apache, MySQL and PHP scripting language installed. The software module constituted of two main parts the front-end and the back-end

4.3.2.1 FRONT-END

The front-end was constructed using HTML. This was the interface which the user interacts with when inputting information to the server or retrieving information from the server. Two main interfaces designed were:

1) An input for the mother’s particulars into the servers. Fields such as Hospital number, admission date and time were auto-completed to ease the process of filling the form.
2) Displaying the measurements recorded by the hardware, simulations as well as the patient’s particulars
An apache hypertext transfer protocol (HTTP) server, a MySQL database management system and PHP scripting language made up the back-end of the server. The apache server was responsible for listening to port 80 for requests from either the user or the hardware. MySQL was used for storage of data inputted from the user and the hardware module. Inputs from the user and hardware modules were processed using PHP which is a server-side scripting language that was used for querying the database as well as processing data to be stored.

4.4 TESTING AND EVALUATION

When developing solutions, it is hoped that it will perform properly in accordance with our vision. However, in practice this is usually not the case as errors and bugs as well as defective hardware influence the resulting outcomes. Thus, the main purpose of evaluation and prototype testing is to find and correct as many errors as possible as well as fixing any failure encountered in the solution. The objectives of the prototype system testing were:

- Ensure system performs as expected
• Test for correctness of result

4.4.1 UNIT TEST

Each of the sub-modules was tested as a separate unit and their outputs studied separately. The hardware sub-modules were checked for correct functionally in comparison to commercially available solution. While the software sub-modules were checked for correctness in data recording and displaying.

4.4.1.1 BLOOD PRESSURE AND HEART RATE

This test dealt with checking whether the results sent to the computer and that displayed by the blood pressure monitor were the same. There being very little information on data being outputted from the blood pressure monitor. Two main steps were taken to establish what kind of data was being outputted by the blood pressure monitor. These included:

• Testing the various standard serial baud rates
• Outputting the data to the serial monitor

For these two tests a simple serial monitor program was written on the Arduino IDE and the output monitored using the Arduino serial monitor. Once the blood pressure monitor started a measurement, data was observed on the Arduino’s serial monitor however the data was unreadable so the baud rate of the blood pressure monitor was adjusted to try and resolve this with no success.
To solve the challenge of this printable data we used minicom a serial monitoring software to dump the content of the serial port onto file. This was achieved using the command ‘\texttt{minicom --device /dev/ttyACM0 --baud 115200 -C capturefile}’ on the Linux terminal with the output being saved into a file named capturefile.

We then used a hexadecimal editor to read the files content. We sought out for patterns that could give us a clue on what the data being transmitted was. This was found at the tail end of the file where the blood pressure monitor outputted two bytes with a value of 253 indicating a successful measurement followed by one byte containing the systolic blood pressure value followed by one byte containing the diastolic blood pressure and another byte containing the value of the heart rate.

\textit{Figure 16: blood pressure monitor test code}
Once this was achieved another program was written to display these readings to the Arduino serial

Figure 17: Binary output from the blood pressure monitor
monitor and the reading on the blood pressure monitor compared for consistency with output on the Arduino serial monitor.

4.4.1.2 TEMPERATURE

This test dealt with checking whether the results from the temperature sensor were valid. After which they were compared to the result of a commercial non-contact infrared thermometer. First the thermometer circuit was connected and the thermometer program loaded with the output being sent to the serial monitor of the Arduino. The result from the commercial thermometer and our prototype were within 0.2 degrees Celsius of each other.

Figure 18: Test result serial output
Figure 19: Prototype temperature test output

Figure 20: Commercial thermometer output
4.4.1.3 URINE

The urine volume measurement sub-module was setup as shown on Figure 10 and an interrupt driven program written to test fluid volume readings. Approximately half a litre of water was placed into a container with a measuring scale. The water was then poured through the flow meter and the serial monitor value compared to the measured value. The outputted value from the controller was approximately the same to that of the measured jar.

![Figure 21: Urine volume sensor serial output](image)

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</thead>
<tbody>
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<tr>
<td>0.520 Litres</td>
<td></td>
</tr>
<tr>
<td>0.0000000000 L/hour</td>
<td></td>
</tr>
<tr>
<td>0.520 Litres</td>
<td></td>
</tr>
<tr>
<td>0.0000000000 L/hour</td>
<td></td>
</tr>
</tbody>
</table>
4.4.1.4 CONTRACTIONS

The contractions measurement circuit was assembled as shown on figure 11 and an interrupt driven program written to test the capture of contraction count as well as handle button debouncing. The Arduino serial monitor output was then compared with the number of times the button was pressed. The resulting output was that for every button press the Arduino registered it and outputted the result to the serial monitor.

4.4.1.5 COMMUNICATION MODULE

The esp8266 Wi-Fi breakout board was connected as show on figure 13. A Wi-Fi router was then configured to connect the server and hardware module. A test program was then loaded on to the microcontroller and connection to the router verified with an internet protocol (IP) number assignment by the router. The output of the test is as shown in the figure below.
4.4.1.6 PATIENT'S PARTICULARS

This test dealt with testing whether the patient particulars were captured correctly.

4.4.1.7 PATIENT RECORD

This test dealt with testing whether the patient records were retrieved and displayed correctly.
4.4.2 INTEGRATION TEST

This test strategy combined all the modules that made-up the prototype. These modules were then tested together to verify that the work together correctly. All the hardware sub-modules were combined and their combine functionality tested.

![Serial output the integrated hardware module](image)

4.5 RESULT AND ANALYSIS

To have a good understanding of the challenges of completing the partograph, 3 previous studies were looked at. Two of the studies were done within a one health facility while the other was done across
9 different health facilities stretching from a health centre to a tertiary hospital. Below is a table showing how partographs were filled.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Completeness (study 1)</th>
<th>Completion (study 2)</th>
<th>Completion (study 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Partograph completion</td>
<td>25.4%</td>
<td>incomplete</td>
<td>&lt;40%</td>
</tr>
<tr>
<td>Admission details</td>
<td>63.3%</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Cervical dilatation</td>
<td>74%</td>
<td>70%-97%</td>
<td>70%-85%</td>
</tr>
<tr>
<td>Foetal descent</td>
<td>54%</td>
<td>minimal</td>
<td></td>
</tr>
<tr>
<td>contractions</td>
<td>49.7%</td>
<td>30%-80%</td>
<td></td>
</tr>
<tr>
<td>Blood pressure</td>
<td>46.9%</td>
<td>minimal</td>
<td>64.8%-71.6%</td>
</tr>
<tr>
<td>Foetal Heart rate</td>
<td>-</td>
<td>53%-90%</td>
<td>72.7%-83%</td>
</tr>
<tr>
<td>Amniotic fluid</td>
<td>-</td>
<td>minimal</td>
<td></td>
</tr>
<tr>
<td>Moulding</td>
<td>-</td>
<td>minimal</td>
<td></td>
</tr>
<tr>
<td>Pulse</td>
<td>-</td>
<td>minimal</td>
<td>62.5%-63.6%</td>
</tr>
<tr>
<td>urinalysis</td>
<td>-</td>
<td>minimal</td>
<td></td>
</tr>
</tbody>
</table>

Table 2: partograph completion rate

From the table it can be noted that completion rate of the partograph is below 40%. Vitals with relatively good completion rate were Foetal heart rate and cervical dilation. Recordings of urinalysis, moulding and amniotic fluid were more or less none existent.

Some of the common reason across the studies for not completing the partograph were poor supervision, no role model, staff shortage and excessive work load.

Result from the prototype hardware module unit test provided good results. The blood pressure and pulse reading were extracted from a commercially available blood pressure monitor that is used in health facilities as well as at home. A feature added to this device was the automatic periodic reading via the hardware control unit. The temperature module based on the DS18B20 chip was also tested.
and its result compared to a commercially available non-contact infrared thermometer. The results from our prototype and the commercial product were mostly the same with occupational variance of about 0.2 degrees Celsius. The flow meter module was tested as well. Once the device was configured a measured amount of water (approximately half a litre) was measured through it and the reading indicated by the module was 520 millilitres which gives a 4% difference. The final measurement module is the contractions this was simulated using a bush button that were registered by our controller for each push. The final hardware module to be tested was the communication module which was tested to see if it would connect to a router and relay data to a server.

The final set of data collected was from 8 volunteers who reviewed the system. The volunteers had backgrounds that included engineering and journalism and have been interacting with medical devices on a regular basis. After making a presentation and demonstration the prototype feedback was obtained using guided question. Only 50% had heard about the partograph, 37.5% were familiar with it, and 25% knew how it was used. On first impressions of the device 6 people participated and give an average score of 80.7%, for usable 5 people participated and gave an average score of 76.8%. The comments give were that the device would assist nurse to fill the graphs and human error would be reduced.
5 CONCLUSION AND RECOMMENDATION

In this chapter a summary of the findings of the research and their implication to maternal care and their stakeholders in this case the nurses and mothers. The chapter sections are; how the study objectives were meet, the contribution of the study, limitations encountered during the study, recommendations and further works to be done.

5.1 HOW OBJECTIVES WERE ACHIEVED

5.1.1 TO INVESTIGATE EXISTING SOLUTION AND HOW THEY ARE USED

In an effort to address this objective, this study sought answers to the objective through literature reviews. The review looked at the stages of labour, what maternal vital signs constituted and the various way the labour process is monitored. Details of this can be found in chapter two of this document.

The chapter describes the labour process as having four stages and the active stage of labour falling between stage one and the end of stage two. The most common tool used to monitor active labour progress is called a partograph. This is a paper-based tool developed by WHO and recommended for use particularly in low resource areas. The tool is divided into three main section that are monitored, these are: Foetal condition, labour progress and maternal conditions.

Other tools developed for labour progress monitoring that employed technology were also looked at, these were the partopen that is a digital partograph solution based on a Dot Position System which uses printed microdots arranged in specific patterns allowing the pen to establish its placement on a form. Another solution looked at was the mlabour which is a mobile application that envelops an electronic partograph into workflows for recording patient details, updating and resolving maternal issues. The final case looked at was the epartogram which is a monitoring and decision support electronic version of the partograph that contains features of both the partograph and the partopen.

5.1.2 TO DEVELOP A PROTOTYPE OF THE DATALOGGER

A prototype solution was developed to meet this objective. The solution composed of two main parts;
the software potion where information was stored and managed and a hardware portion on which all the sensory systems were housed. The sensors used where:

- A blood pressure and heart rate monitor for measuring the mother’s systolic blood pressure, diastolic blood pressure and heartbeat
- A temperature sensor for the measurement of the mother’s temperature
- A flow meter for measuring urine volume
- A push button for counting contractions

A communication module was also included and it was responsible for transmission of data to the server. At the heart of the hardware was a control unit responsible for time keeping, measurement and interpretation of signals from the sensors as well as transmission of the data to the server.

5.1.3 EVALUATE THE SYSTEM

This study pursued a look at the possibility of automating the measurement and logging of maternal vital signs during the active stage of labour and their retrieval to assist in the completion of the partograph.

Testing of the solution was mainly confined to laboratory experiments and feedback from colleagues. Feedback from the test indicated that the solution faithfully monitored and recorded the vitals as well as gave an accurate reading of the vitals measured. From the group discussion the idea was thought to be useful and could assist nurses in plotting their graphs, it would also reduce the occurrence of human errors that occur during measurements and recording.

5.2 CONTRIBUTION OF THE STUDY

This research sought to develop a digital information technology solution to assist nurses and midwives record labour progress during the active stage of labour. The prototype created a platform in which some of the periodic vital signs measured during labour were automated: these included blood pressure, heart rate, temperature, contractions, and urine volume. These signs were periodically measured and recorded to a server from where they could later be retrieved to assist in completion of the partograph. Hence removing the need of nurses having to generate random data to complete a partograph as well as enabling them attend to mothers more effectively.
5.3 STUDY LIMITATIONS

Various limitations were encountered within this study. Design of a data logging solution proved to be a technical challenge as material resources for use in this project proved to be unavailable resulting in sourcing of some of the material resources from outside the country which in-turn took time to be delivered. This resulted in a longer duration would be required to identify and source for needed material as well as testing their functionality.

Another limitation had to do with the medical nature of this study. Acquiring ethical approval to conduct research in any health facility was required. This process takes between one and two months at best. To go around this secondary data from other research works was used as a foundation of some of the findings as well as feedback from persons working on medical devices pertaining to maternal care.

For a good evaluation of the solution clinical trials would give a good indicator of how well the solution performs in the real world in the hands of actual nurses and mothers. However, this would take a few months to conduct hence not possible within the time period of the project as well as need ethical approval as mentioned above.

5.4 RECOMMENDATION AND FURTHER WORK

This research focused mainly on monitoring of maternal vital signs. The decision to focus on digitization of maternal vital signs was arrived at due to limitation of locally available technologies, time resource, financial constraints and technical complexity of measuring and recording of some of the signs such as automation of visual inspection, drug administration where applicable and chemical analysis. Hence there is still a need to create a digital solution that is capable of monitoring, measuring and recording more if not all the vitals on the partograph.

Sharing of information among the health facilities is also another point of note This would enable better service provision to mothers in referral cases as the hospital the mother has been referred to will easily be able to conclude on the exact state the mother and baby are in without having to perform their own independent test and analysis.

Clinical trials of the prototype would also need to be performed to test the performance of the prototype under real world conditions. This would shed more light on limitations and advantages of
its functionality by the nurses, unforeseen failure points and its capacity to perform in a real-world test compared to laboratory testing.
REFERENCES


APPENDICES

APPENDIX A: CONFIDENTIALITY AGREEMENT FORM

I Kennedy Abwao Ogendo, Reg. No P51/65174/2013, a bona fide student of the University of Nairobi, School of Computing and Informatics agree that as the principal investigator executing this research exercise, I will ensure that all information captured and handled in whatever form in the process of carrying out this research exercise will be kept confidential and will not at any point in time; during and/or after the research exercise be disclosed to any other person or used for any other purpose other than the intended research.

Signature............................................................

Date....................................................................
APPENDIX B: INFORMED CONSENT NOTE

Research Title
A DATA LOGGER FOR MONITORING MATERNAL VITAL SIGNS DURING LABOUR

Introduction: Background

I Kennedy Abwao Ogendo, a MSc student at the University of Nairobi School of Computing and Informatics intend to carry out a study that aims at evaluating interventions that can reduce maternal mortality and the outcomes resulting from these interventions.

Research Procedure Brief

The research procedure and process involve interacting with healthcare workers who in one way or another interact with expectant mothers or mothers in labour. Data will be collected from the respondent using questionnaires. We will analyse the data collected in an effort to understand the interventions that can reduce maternal mortality and the outcomes resulting from these interventions.

Confidentiality Issues

All information captured will have no respondent’s identifier (Huduma Number, ID Number, Name, Passport Number, Birth certificate Number etc.). After collection, all data and information will be in the custody of the principal researcher for safe and secure keeping.

Respondents Voluntary Participation

Each respondent will voluntarily participate in the study after a briefing on what the study is all about. They will be required to read and understand the informed consent note and seek all clarification where necessary. Each respondent will be allowed time to make an independent decision on whether or not they would like to participate in the study. They will then be required to sign a consent form as evidence of their voluntary participation in the research.
Respondents Eligibility to Participate

Respondents of this study will be required to fulfil the following requirements:

- Should be one who interact with expectant mothers or mothers in labour
- Should voluntarily agree to participate in the study

Motivation and Drive

Our motivation for this study is to establish and understand the outcome of interventions with respect to maternal healthcare. This will provide a knowledge base that can help in the creation of solutions that will mitigate mortality caused by labour related factors.
APPENDIX C: CONSENT FORM

I acknowledge that I have read and understood Informed Consent Note concerning this research and that I have also sought all necessary clarification concerning my participation as a respondent in the research. By appending my signature, I voluntarily agree to participate in this particular study.

Names...........................................................................................................

Signature ....................................................................................................... 

For any further information or clarification about this research, please contact:

Kennedy Abwao Ogendo
C/o University of Nairobi,
School of Computing and Informatics
P.O. Box 30197 – 00100
Nairobi, Kenya
Tel: +254 -721 587 688

If you (respondent) have any concern or complaint relating to the conduct of this research, kindly contact:

The Chairperson
KNH-UoN Ethics Review Committee
P.O. Box 19676 - 00202
Nairobi, Kenya
Tel: 020-272 6300-9 ext. 44355
APPENDIX D: QUESTIONNAIRE
PART A: BRIEF ON THE RESEARCH

This questionnaire is part of a research by Kennedy Abwao Ogendo towards a Master of Science Degree at the University of Nairobi, School of Computing and Informatics. The study seeks to review and understand the interventions that can reduce maternal mortality. Possible solution for mitigating some of the healthcare challenges associated with maternal care will then be designed based on this data.

PART B: IMPORTANT NOTE TO THE RESPONDENT

NO respondent’s personal identification information should be filled anywhere in this questionnaire. These will include but not limited to those listed below

- Huduma Number
- National Identity Card Details (Number and Names)
- Passport Details (Number and Names)
- Birth Certificate Number
- Hospital Records, employee number

PART C: WORKING CONDITIONS

1. How many colleagues do you work with in a shift? ______________________________
2. How long is a shift? __________________________________________________
3. How many mothers do you have to attend to in a shift? (less than, between, more than)
4. How many shifts does the labour process take? (1 shift, 2 shifts, more than 2 shifts)

PART D: SKILLS AND MONITORING TOOL

5. On a scale of 1 – 10 (1=not available, 10=ready available) how available are partographs for use to monitor labour? __________________
6. On a scale of 1 – 10 (1=easy, 10=hard) how available are partographs for use to monitor labour? __________
7. How often do you get training on the use of the partograph? (never, occasionally, frequently)
PART E: TOOL COMPLETION LEVEL

8. Please fill the table below with respect to the partograph

<table>
<thead>
<tr>
<th></th>
<th>On a scale of 1-10 how often are these reading taken during labour progress? (1=never, 10=always)</th>
<th>How is the vital recorded? (visual inspection, using a tool (please mention tool) or both)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contraction</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cervical dilation &amp; effacement</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Descent of head</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urinalysis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mother’s blood Pressure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mother’s temperature</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mother’s pulse</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Respiratory</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amniotic fluid moulding</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Foetus pulse rate</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

9. How long does it take to complete one entry on the partograph? ___________________

10. How much time do you think you need to measure and record all the data required in the partograph for all the mothers you need to attend to? ____________________________

PART F: INTER-FACILITY COMMUNICATION

11. Do you get referrals? (YES / NO)

   If yes
   1. What state are the mothers in on a scale of 1-10 (1=normal, 10=critical)? _____
   2. Do the mothers come accompanied with a partograph in a scale of 1-10? (1=never, 10=always) _______________
      1. if yes, how complete are they on a scale of 1-10? (1=blank, 10=completed)
         __________________________________________________________
      2. If yes, how helpful is it in giving the state of the mother? (1=not at all, 10=very
12. Is there anything important you think I may have missed?

__________________________________________________________________________

__________________________________________________________________________

__________________________________________________________________________

CONTACT:
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Nairobi, Kenya.