Monitoring intrapartum fetal heart rates by mothers in labour in two public hospitals: an initiative to improve maternal and neonatal healthcare in Liberia

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Abstract

Background: In low-resource settings with few health workers, Fetal Heart Rate (FHR) monitoring in labour can be inconsistent and unreliable. An initiative to improve fetal monitoring was implemented in two public hospitals in rural Liberia; the country with the second lowest number of midwives and nurses in the world (1.007 per 10,000 of the population). The initiative assessed the feasibility of educating women in labour to monitor their FHR and alert a midwife of changes detected. Methods and Interventions: 474 women admitted in labour without obstetric complications were approached. 461 consented to participate (97%) and 13 declined. Those consenting were trained to monitor their FHR using a sonicaid for approximately one minute immediately following the end of every uterine contraction and to inform a midwife of changes. If changes were confirmed, standard clinical interventions for fetal distress (lateral tilt, intravenous fluids and oxygen) were undertaken and, when appropriate, accelerated delivery by vacuum or Caesarean section. Participants provided views on their experiences; subsequently categorized into themes. Neonatal outcomes regarding survival, need for resuscitation, presence of birth asphyxia, and treatment were recorded. Results: 461 out of 474 women gave consent, of whom 425 (92%) completed the monitoring themselves. 387 of 400 women who gave comments, reported positive and 13 negative experiences. 28 participants reported FHR changes, confirmed in 26 cases with meconium stained liquor in 17. Fetal death was identified on admission during training in one mother. 13 neonates required resuscitation, with 10 admitted to the neonatal unit. One developed temporary seizures suggesting birth asphyxia. All 26 neonates were discharged home apparently well. In 2 mothers, previously unrecognized obstetric complications (cord prolapse and Bandl’s ring with obstructed labour) accompanied FHR changes. Resuscitation was needed in 8 neonates without identified FHR changes including one of birth weight 1.3 Kg who
could not be resuscitated. There were no intrapartum stillbirths in participants. Conclusions: Women in labour were able to monitor and detect changes in their FHR. Most found the experience beneficial. The absence of intrapartum stillbirths after admission and the low rate of poor neonatal outcomes are promising and warrant further investigation.

Background

The need to improve identification and management of intrapartum placental insufficiency and fetal distress is particularly relevant in low-income countries, such as Liberia. Yet it is in such settings that the methods and means to do so are severely limited by the lack of health workers and by material poverty. Challenges include the lack of suitably skilled birth attendants, particularly in rural settings; the high workload with each attendant often having to care for several women in active labour, especially during the night; and the lack of resources to fund, maintain and interpret fetal monitoring devices. As quoted by The Lancet Every Newborn Study group: “Sensitive, specific, and simpler methods for detection of fetal compromise during labour could have a major effect on intrapartum stillbirths and early neonatal deaths, as long as linked with emergency obstetric care” [1].

The scale of the shortage of suitably trained health workers, such as nurses and midwives to undertake any fetal monitoring is highlighted in Table 1, which shows the numbers of nurses and midwives per 10,000 of the population as determined by WHO in 2019 in its Global Health Observatory Data Repository [2]. 17 out of 194 countries had < 5.0 midwives and nurses per 10,000 of the population and 15 out of 194 countries had > 100 midwives and nurses per 10,000 of the population

Table 1 near here. Lowest and highest numbers of nurses and midwives per 10,000 of the population in 194 countries in the world.
The two countries with the lowest numbers of midwives and nurses currently recorded by WHO are Somalia and Liberia. Both countries have experienced recent armed conflict. Between 2014-2015, Liberia also experienced an Ebola outbreak, which killed many healthcare staff. Unfortunately, since recovery from Ebola, the international community’s statements that they would improve the public health infrastructure of Liberia have not materialized [3].

Table 1 also shows that high-income countries have more than 100 times the numbers of midwives and nurses than Liberia, available to attend pregnant women and their babies. Analysis of the WHO Global Health Observatory Data Repository [2], on the numbers of doctors, shows that out of 194 countries in the world, Liberia has the 4th smallest number of doctors (0.373 doctors for 10,000 of the population).

Moreover, as a result of increasing inflation in Liberia, (7.7% in 2009 to 22.3% in 2019) [4] and worsening poverty, the employment of midwives in the public hospitals has reduced, their salaries have decreased, and sometimes, they may not be paid for several consecutive months. Subsequently, their morale has fallen, and it is rare for partographs to be completed according to WHO recommendations. Although the Ministry of Health has made great efforts to educate midwives to international standards as recommended by WHO [5], the lack of funds available to the national health service facilities, frequent absence of essential drugs, medical and surgical supplies, together with a lack of international funding, makes providing high quality health care difficult to achieve, especially in rural areas where attracting health workers of all levels is difficult. Labour wards in rural Liberian hospitals are frequently over-crowded with patients and it is rare for any mother to in labour to be accompanied by a relative or for them to receive the levels of attention and care, both to themselves and to their unborn babies, which are essential to prevent and / or treat intrapartum complications.
A study conducted in a Liberian referral hospital showed that out of 1656 deliveries in one year (2010), there were 196 perinatal deaths; 143 classified as stillbirth and 53 classified as early neonatal death. [6].

The majority of stillbirths (56.6%) presented as antenatal stillbirths with no fetal heart sounds documented on admission. Thirty-seven (25.9%) had documented fetal heart rates upon admission with the stillbirth occurring during the intrapartum period. There was no documentation of presence or absence of fetal heart rates in 25 of the stillbirth records (17.5%). Thirty-one percent of cases had no maternal or obstetrical diagnosis recorded in the chart when a stillbirth occurred. Of the 53 early neonatal deaths, 47.2% occurred on the first day of life. The largest single contributor to early neonatal death in the sample was birth asphyxia identified by poor Apgar scores (< 4 at 5 minutes). Seventeen of the 53 early neonatal deaths (32%) were due to birth asphyxia.

After our health improvement initiative / feasibility study began in July 2017, WHO in February 2018 produced an updated guideline concerning intermittent FHR auscultation during labour using either a Doppler ultrasound device or a Pinard fetal stethoscope. [7] This WHO expert group reported that monitoring of the fetal heart rate during labour was inadequate in many low- and middle-income country (LMIC) settings, and that this problem needed to be strongly addressed through quality improvement initiatives in these settings. In their review of qualitative studies exploring women’s experiences of labour and childbirth, WHO suggested that women would prefer a more hands-on, woman-centred approach to care and were likely to favour any technique that allows for this approach, [8] WHO also pointed out that Doppler monitoring allows a woman to hear the fetal heartbeat, which provides reassurance and could add to its appeal over Pinard auscultation. Their guideline also referred to WHO’s 2015 State of inequality report [9], indicating that women who are poor, least educated, and residing in rural and remote
areas have lower access to healthcare intervention coverage than more advantaged women. The new guideline also emphasised studies reporting that adequate monitoring of labour progress is often lacking in such settings, and that the FHR may only rarely be auscultated [10-11]

Such rare use and undertaking of FHR monitoring represented the situation in rural public hospitals in Liberia in 2017, which led to the design and implementation of this present initiative.

Globally, intrapartum-related complications are reported to cause an annual 1.2 million stillbirths, 700,000 term newborn deaths, and an estimated 1.2 million newborn babies developing neonatal encephalopathy (birth asphyxia) with 233,000 survivors developing moderate or severe neurodevelopmental impairment [12]. The countries with the highest stillbirth and neonatal mortality rates are in Sub-Saharan Africa [1], where between 25.1 and 34.2 stillbirths occurring for every 1,000 births, with an estimated 51% of these deaths happening intrapartum [13]. For example, in a rural hospital in Tanzania, the stillbirth rate was 27/1000 live births, with 16/1000 occurring intrapartum [14] and 27% of deaths in the first 6 days of life related to intrapartum causes.

Compared with well-resourced countries, intrapartum-related neonatal mortality rates are 25-fold higher and intrapartum stillbirth rates up to 50 times higher in the lowest-income countries [15] where rehabilitation services for children with neuro-developmental impairments are poor or absent.

The integration of obstetric and neonatal care to manage complications for both mother and fetus during labour [16] is critically important with task-sharing being particularly relevant in low resource settings. To the best of our knowledge, mothers have not previously participated in FHR monitoring during labour. Their contribution (as a form of task-sharing) could be valuable for the wellbeing of both themselves and their babies and
could be of assistance to overworked health care professionals who are too busy to undertake regular fetal monitoring. WHO recommendations for intrapartum fetal monitoring by midwives may be possible to achieve in well-resourced countries (100-180 nurses and midwives per 10,000 population [2]) but may be aspirational in poorly resourced countries, such as Liberia, which has 1.007 nurses and midwives per 10,000 population [2].) As part of a maternal and neonatal health care improvement initiative, we assessed whether maternal participation in monitoring the FHR during labour is a potentially feasible, effective and sustainable approach to improving intrapartum and neonatal survival in situations with few and overworked skilled birth attendants.

Methods

Specific Aims and Methods

We have used the revised standards for quality improvement reporting excellence (SQUIRE 2.0) to report this initiative [17].

Given the major shortage of midwives in Liberia, the first aim of this initiative, was to assess the feasibility of educating women in labour to assess their FHRs with a portable doppler monitor and to alert the midwife if they detected changes in the FHR which might indicate fetal distress.

The second aim of this initiative was to facilitate system changes to educate and enable the attending midwife, who would likely be caring for several individual women in labour, to initiate an immediate remedial course of action should she/he be alerted by a woman in labour who detected changes in her FHR. The questions posed, and the measured outcomes of this intervention, are shown in Table 2.

Table 2 (near here) Questions to be addressed by this initiative.

Setting

In Liberia, over the past 6 years, a partnership between the Ministry of Health and Social
Welfare (MOHSW), The World Health Organization (WHO), The United Nations Population Fund (UNFPA), the Liberian Board for Nursing and Midwifery (LBNM) and the registered charity (not-for profit organization), Maternal and Childhealth Advocacy International (known as MCAI), has been established to train senior birth attendants (19 midwives and 2 physician assistants) in advanced obstetrics to become obstetric clinicians, able to undertake emergency clinical procedures to expedite the delivery of a distressed fetus, such as vacuum delivery and Caesarean section [18]. Currently, ten obstetric clinicians have completed training and 10 more are currently in their final year of training.

More recently, the partnership initiated a new programme to improve neonatal resuscitation and hospital neonatal care by training selected nurses and midwives to become advanced neonatal nurse practitioners (neonatal clinicians) and is now based in both of the two hospitals involved.

The fetal monitoring initiative took place in CB Dunbar Hospital and CH Rennie Hospital. The main training hospital for both obstetric clinicians and advanced neonatal nurse practitioners (neonatal clinicians) is CB Dunbar Maternity Hospital in Bong County. During this initiative, there was one fully qualified and licensed obstetric clinician, 4 trainee obstetric clinicians and 9 trainee neonatal clinician practitioners at CB Dunbar Hospital. There were 5 trainee obstetric clinicians at CH Rennie Hospital but no neonatal clinicians at the time of this feasibility study.

*Management committee for this health care improvement initiative*

This initiative represented an integrated project between obstetrics and neonatology. Both qualified and trainee obstetric clinicians and trainee neonatal clinicians and their trainers were involved.

The management committee oversaw the day to day running of the initiative and was responsible for the data analysis. If there were any problems identified, in particular
involving the need for improving the recording methods, the management committee
would make the necessary changes to the initiative as appropriate.

As members of the management committee, two trainee Obstetric Clinicians (KP and NJ),
and the international advanced neonatal nurse practitioner (AK) managing the recently
established neonatal intensive care unit, led the day to day running of the fetal
monitoring component of this initiative.

Participants

Over a 15-month period (from 31 July 2017 to 24 October 2018), 474 women admitted in
the active first stage or second stages of labour, who were not experiencing any
complications such as haemorrhage, severe pre-eclampsia or obstructed labour, were
invited to participate. Before participation, each mother was asked to give her informed
consent by a trainee or fully trained obstetric clinician. Initially (given a female literacy
rate from UNICEF data 2011-2016 [19] of those aged 15 to 24 years of approximately
37%) this consent was requested verbally and recorded and later (after 27\textsuperscript{th} April 2018) a
dedicated consent form was signed (either by fingerprint or in writing). The consent form
(part of the Additional File) was read out and discussed with each woman before she was
enrolled in the study. Only women or adolescent girls (those potential participants aged
under 18 years) who fully understood what they were being asked to undertake were
recruited. The obstetric clinicians who gained consent were skilled health workers with
years of experience in managing pregnant women in Liberian public hospitals. All had
undergone a medical ethics and professional standards course as part of their training.
Therefore, these health workers were well placed to ascertain if each woman who was
invited to participate understood what was required. Each woman could change her mind
at any time and her wishes were always respected.

Plan of investigation
Between contractions, each participating woman was shown and educated by an obstetric clinician in how to use a re-chargeable, battery operated, fetal doppler heart sound monitoring device (a sonicaid), including the best position to place the probe on the mother’s abdomen. This training usually lasted between 10 and 15 minutes. The mother was also educated in what was a normal heart rate and what was slow or fast, by tapping out a rhythm. Mothers were asked to monitor for approximately a 60 second period immediately following the end of every contraction. However, it wasn’t, in our opinion, of crucial importance or appropriate to insist that participating women monitor for exactly 60 seconds, which may be difficult. In addition, it is/was the beginning of the 60 second period that we considered of most clinical relevance to detecting changes in the FHR that might indicate fetal distress.

Having determined what was a normal rate, the mother was then asked to identify and immediately inform a midwife of any changes in the FHR that she detected. In the hospitals of this initiative, there is no buzzer or bell alert and so if a participating woman thought that she had detected a change in FHR, she called out for a midwife or obstetric clinician to come and confirm possible changes. As the labour wards in both hospitals were small, this system worked well. Sometimes, the mothers reported actual heart rates (if literate) but mostly identified a fall or increase in heart rates without counting the numbers involved. The midwives and obstetric clinicians who responded to the mother, recorded and reported the actual heart rates. An Additional File contains the form that was used for the recording process which was completed as soon as possible after delivery.

The Additional files also contain two tables summarizing the clinical data collected on all mothers approached for consent, together with the comments made on the monitoring process by the mother, either written directly or transcribed for illiterate mothers by the
obstetric clinician completing the form.

In all participating women, the partograph was supposed to be completed as normal by a midwife on the labour ward every 30 minutes during the first stage and every 5 minutes during the second stage of labour [20,21]. However, in reality the fetal section of the partograph was inconsistent and rarely completed due to time pressures on the small number of midwives.

If a participating woman decided that she could not continue to monitor the FHR, for whatever reason, her wishes were respected and documented. If a woman's condition during labour made it difficult for her to continue making recordings, a midwife or obstetric clinician would, if possible and if time allowed, assist or take over the monitoring from her with the woman’s permission.

If a woman declined to undertake the monitoring herself but still wanted her fetus to be monitored, with the mother’s permission, the monitoring could be done by a midwife or by an obstetric clinician, provided they had time available given their heavy workloads.

**Actions by midwives and/or obstetric clinicians**

The midwives at the two hospitals were informed of the plans and scope of the initiative by the management committee team and educated in their role and responsibilities regarding the initiative, especially that they must, if possible, respond immediately when alerted by the participating women to a change in FHR and to examine the woman. If the FHR was potentially of concern, the midwife would then immediately notify the obstetric clinician and/or doctor on duty who would assist the midwife in the management of labour until the FHR had either recovered or the baby was safely delivered (as per the birth asphyxia prevention protocol: see below). This remedial protocol was already in place on the maternity units should potential fetal distress be detected. However, unfortunately, FHR monitoring as part of the partograph was rarely undertaken prior to this initiative for
reasons outlined in the earlier “Background” to this project. At CB Dunbar Hospital, the neonatal clinician on duty would also be asked to be present at the delivery to resuscitate the neonate if necessary.

*The Birth Asphyxia Prevention Protocol*

1. If a participating woman in labour detected a possible FHR change (bradycardia or tachycardia) she immediately notified an available midwife.

2. The midwife checked the FHR and the mother’s condition.

3. If a bradycardia (<120 beats/minute) or tachycardia (> 160 beats/minute) were present, the midwife immediately notified the obstetric clinician or doctor on duty.

4. If the FHR was normal, the midwife immediately began a period of continuous fetal monitoring including the time up to the next contraction. She / he also listened to the FHR during the whole of the next contraction and immediately following that contraction.

5. If a suspicious FHR was detected during the one minute following the next contraction, the midwife took the following actions:
   (i) ensured the mother was not lying flat on her back by providing left lateral tilt,
   (ii) ensured that the obstetric clinician and/or doctor on duty was present
   (iii) provided additional inspired oxygen (when available), secured an intravenous cannula and give a bolus of either 0.9% saline or Ringer Lactate solution and where there was suspicion of ketosis, added a bolus of 50% glucose to the IV infusion [21]

6. If there was evidence of fetal distress (late decelerations, persistent bradycardia, persistent tachycardia, meconium stained liquor) then the mother was assessed by the obstetric clinician or doctor to manage any maternal obstetric problem that could
be responsible for the fetal bradycardia/tachycardia and assess for urgent immediate
delivery as follows:

(i) If the cervix was fully dilated and there were no contraindications to
instrumental delivery, a vacuum delivery was undertaken.

(ii) if the cervix was not fully dilated, then the woman was prepared for an
emergency Caesarean section (CS) and taken to the operating theatre where she was re-
examined. If the fetus was still alive and the cervix still not fully dilated, a CS was
performed. An abdominal ultrasound scan was often helpful at this time, not only to check
whether the fetus was still alive, but also related to other clinically relevant issues, such
as to determine the lie and position of the fetus and the position of the placenta.

7) Under either circumstance outlined in 6 above, a neonatal practitioner or midwife
experienced in neonatal resuscitation, was immediately made available for when the baby
is delivered. She/he would ensure that the equipment needed for resuscitation was
available and functioning.

The outcome for the mother and fetus/baby.

The clinical condition of the baby at birth, the need for neonatal resuscitation, admission
to the neonatal unit and any signs of subsequent birth asphyxia (Hypoxic Ischaemic
Encephalopathy: HIE) were documented. A clinical summary of the pregnancy and delivery
was documented.

The views of the mothers on the fetal monitoring process.

From 23rd October 2017, through written or verbal comments (the latter transcribed by
the midwife or obstetric clinician), mothers were asked for their views on the monitoring
process.

153 out of 154 participants who had consented at CH Rennie hospital (where the project
started on 21st March 2018) were approached for comment (on admission one potential participant’s fetus was found to be dead during her training in monitoring).

This request for comments was not in place for the 33 of the consented mothers, who had enrolled before 23rd September 2017 at CB Dunbar Hospital.

Two authors (DS and RM), categorized the comments retrospectively by first agreeing the common themes after reading all of the comments provided by the participating women. Any discrepancies between DS and RM were discussed and agreed by consensus.

The comments in Table 3 were selected as being representative of the basic themes, as subjectively agreed by the two authors RM and DS. The full set of comments from all participants are available in the Supplementary Information (Additional files).

Data analysis

MCAI was responsible for the data analysis and undertook a descriptive analysis.

The lead obstetric clinician in each hospital (KP and NJ) was responsible for charging the batteries and for ensuring gel, consent, and monitoring forms were constantly available. Forms were scanned and sent to the UK by MCAI programme and finance managers in Liberia.

Figure 1 near here: A mother checking her unborn baby’s heart rate.

Legend to Figure 1. One of the mothers at CB Dunbar Hospital checking her unborn baby’s heart tones during the first stage of her labour immediately following a uterine contraction using a sonicaid (fetal doppler battery operated machine). Written consent to use this picture in this publication was obtained.

Results

Figure 2 near here. Flow chart showing the evolution of the project

Legend to Figure 2. A Flow chart which describes the timelines of the various
interventions undertaken up to the completion of this manuscript. The project meanwhile continues in both hospitals.

Consent and monitoring  (see flow chart Figure 3)
Figure 3 near here. Flow chart showing consent and comments for this initiative
Over 15 months from 31 July 2017 until 24 October 2018, 474 women (157 from CH Rennie and 317 from CB Dunbar Hospitals) admitted in the first or second stages of labour and without obstetric complications were approached. 461 gave their informed consent to participate in the FHR monitoring (97%) and 13 declined.

CH Rennie hospital data
At CH Rennie Hospital, data were collected from 21 March 2018 until 24 October 2018. 157 mothers were approached for consent. Three declined [case number 6 (too much pain); case number 54 (no reason); and case number 75 (no reason). In the 3 women who declined, only fetal monitoring as part of the partograph, continued to be undertaken. Three (case numbers CHR 4, 10 and 153) had consented but, after monitoring for 12, 8 and 36 contractions respectively, stopped monitoring because of pain and tiredness. Fetal death was identified on admission during training in the use of the sonicaid in one mother (case CHR 50 in Table 4: see below).

CB Dunbar hospital data
At CB Dunbar Hospital, data were collected from 31 July 2017 until 23 October 2018. 317 mothers were approached for consent. 307 gave their informed consent and 10 declined. In 4 of these cases, where the mother said her pain was too much to allow her to do it, with the woman’s permission, the obstetric clinician and/or midwife continued to undertake the monitoring with the sonicaid. In one of these latter 4 cases (CBD 251) a student midwife undertook all of the monitoring and identified a change in FHR (see Table
4). In the other 6 cases, no FHR monitoring was undertaken except as part of the partograph.

Out of the 307 mothers who consented, there were 7 cases where the mothers initially declined but then changed their minds and undertook monitoring for the rest of their labour. In one of these cases, the mother shared the monitoring with her midwife and obstetric clinician.

In an additional 7 cases, the mother consented but subsequently stopped monitoring (1 through tiredness, 1 because of pain, 4 with no reasons given and 1 because she was worried about delivering on time).

In an additional 20 cases where the mother consented but then stopped monitoring, a midwife or obstetric clinician took over (19 cases) until delivery or shared the monitoring with the mother (1 case). Of these 20 cases, 15 gave pain and 5 gave tiredness or weakness as the reasons.

425 of 461 (92%) participating mothers were able to complete the monitoring themselves (151 from CH Rennie Hospital and 274 from CB Dunbar Hospital).

Maternal age

Maternal age was available for 416 of 461 participants. 48 (12%) were aged under 18 years; 22 mothers aged 17; 17 aged 16; 5 aged 15; 3 aged 14; and one aged 13 years.

Maternal experiences (including evidence regarding the consequences of the lack of any pain control, which was not available during labour in any of the public hospitals in Liberia).

400 participants provided written or verbal (transcribed) comments on their experiences of the monitoring.

387 mothers found listening to their unborn baby a positive experience expressing one or more of the following words or phrases: alright, not bad, good, fine, helpful, loved or liked.
it, happy, comfortable, gives me joy, or other positive comments such as “Thank you”. A selection of these comments is shown in Table 3, including some from mothers who identified changes in FHR. The complete set of comments from all participants are available in the supplementary information (additional files).

13 participants reported only negative comments: 5 reported weakness or tiredness (including feeling nauseated in one case), 6 reported pain (which in 3 interfered with the monitoring), 1 said it was not easy and 1 said it was bad.

Table 3 (near here) Selected maternal comments from mothers at both Hospitals (additional maternal comments following changes in fetal heart rate are reported in Tables 4 and 6 below).

Out of the 387 providing positive comments, 86 also reported how much they were affected by pain or severe pain. This pain interfered with their ability to undertake the monitoring in 52 of the 86 (60%).

Within the 387 with positive comments, 44 women also reported discomfort, 5 reported tiredness and weakness and 4 reported difficulties applying the sonicaid.

Three mothers said that they would hope that monitoring would be available during their future pregnancies. 6 said they would return to the hospitals to deliver as a result of the project. An additional 6 mothers said they would hope that monitoring will continue to be available so that other mothers can benefit and 5 said they would encourage other mothers to participate. 9 participants said monitoring helped to cope with labour pain.

Technical and administrative problems identified

As the forms were to be completed by busy midwives and obstetric clinicians, not dedicated researchers, forms were designed to record the most relevant information for the purposes of the study objectives rather than detailed information which, although useful, was not necessary for the purposes of this initiative and could have taken the
health workers away from their clinical work.

The initial design of the tick sheet documenting each contraction monitored did not always allow enough space to record every contraction and obtaining extension sheets was sometimes a logistic problem. To minimise the workload of the scarce midwifery workforce, forms were also re-designed to record clinical data that were appropriate but not excessive given time restraints.

Due to a communication problem, 57 mothers at CB Dunbar Hospital (between 19 Feb 2018 and 8 May 2018) incorrectly monitored their FHR every 30 minutes (similar to the partograph). However, unlike the partograph, monitoring was always undertaken for approximately 60 seconds immediately following the nearest contraction to each 30-minute window. We did not identify any clinical differences or differences in satisfaction between the 57 women who monitored the fetal heart every 30 minutes and the women who monitored after every contraction.

Birth/delivery data

In 461 participants, there were 33 caesarean sections (7.2%) including 14 with FHR changes. There were 20 vacuum deliveries (4.3%) including 8 with FHR changes, (the latter included a mother with a stillborn baby identified during the training in the use of the sonicaid, confirmed by ultrasound scan).

Clinical information and outcomes for participants where changes in FHR were identified.

Table 4 near here. Clinical information and outcomes where FHR changes were identified by monitoring.

Table 4 describes the clinical information and outcomes relating to identified FHR changes. Changes in FHR were reported in 28 of 461 participants (6.1%), which in two cases were not confirmed by the attending midwife, giving 26 confirmed cases (5.6%). In 23 of the 26 confirmed cases, the FHR decreased and in 3 the FHR increased. Two
changes related to unrecognized obstetric complications, with one mother found to have Bandl’s ring with obstructed labour and the other, cord prolapse. 

All 26 neonates with changes identified in FHR survived, including 13 requiring resuscitation at birth. Twelve neonates were admitted to the neonatal unit. One neonate developed birth asphyxia/HIE (case CHR 99) but the immediate clinical outcomes of the 25 other neonates were good and all 26 were discharged home apparently well. One of the 26 changes in FHR was identified by a midwife who had taken-over monitoring from a mother who became too tired to continue. In a second case, the mother had declined to undertake monitoring herself but had consented to the monitoring being undertaken by a student midwife.

In 17 of the 26 with confirmed FHR changes plus 1 intrauterine fetal death detected on admission, there was accompanying meconium-stained liquor. 13 of the 26 (50%) neonates with prior FHR changes had low Apgar scores and needed resuscitation. There were no deaths following resuscitation. One baby (case CHR 99) had convulsions managed with Phenobarbital, recovered and was feeding normally at discharge home aged 7 days. None of the other 26 neonates developed birth asphyxia (also known as Hypoxic Ischaemic encephalopathy-HIE). However, long-term infant follow-up was not undertaken.

8 of the 26 neonates with confirmed FHR changes plus 1 with intrauterine fetal death (30%) were born by vacuum, 14 by Caesarean Section (CS) (54%), and 6 (23%) by vaginal delivery. In one case (CBD 272), CS followed a failed vacuum delivery.

Clinical information and outcomes in 3 newborn infants needing resuscitation at birth where mothers had declined to participate in FHR monitoring.

TABLE 5 (near here). Clinical information and outcomes in newborn infants needing resuscitation at birth where mothers had declined to participate in FHR monitoring.
In one such case, the baby was born with Apgar scores of 2 at 1 minute and 3 at 5 minutes and, despite resuscitation, died of HIE in the neonatal unit aged 2 days. In 2 other cases, there were low Apgar scores at 1 and 5 minutes (4 and 6; and 2 and 6) and the babies needed resuscitation. Both were admitted to the neonatal unit. One responded well to resuscitation with no evidence of HIE and was discharged home well. The other died aged 3 days from birth asphyxia/HIE.

Clinical information and outcomes in 8 neonates needing resuscitation where no FHR changes had been identified

TABLE 6 (near here). Clinical information and outcomes in newborn infants needing resuscitation at birth where monitoring had not identified any FHR changes.

In addition to the neonates requiring resuscitation where FHR changes had been detected, 8 other neonates without detected changes in the FHR required resuscitation (Table 6). One (case CBD 238) was preterm/low-birth weight and died at birth. None of the remaining 7 developed birth asphyxia/HIE.

In one neonate (Case number CBD 224), it was unclear who had undertaken the monitoring and for how long. A vacuum delivery was undertaken for failure to push, Apgar scores were 5 and 8 and the baby required 10 minutes of bag and mask ventilation. He was discharged home aged 7 days well. No evidence of birth asphyxia/HIE was evident on clinical assessment.

Three cases were born following vacuum delivery and 5 cases by vaginal delivery, including in one mother where an episiotomy was undertaken to expedite delivery (case CBD 240).

Three of the neonates had been monitored in utero only every 30 minutes in a temporary deviation to the protocol because of a communication problem with one of the trainee obstetric clinicians but none developed birth asphyxia/HIE.
Costs associated with collecting data

The costs of the project were low. The fetal doppler monitors (Sonicaid: 12 in total) were USD 40 each. Rechargeable AA batteries were used. Additional costs included paper and printing for the consent, data collection and monitoring forms including the internet costs of scanning and sending them to MCAI for analysis and KY jelly (or locally available clear hair gel) for interfacing the ultrasound probe with the abdomen: commercial ultrasound gel was too expensive.

Missing data

Because of problems with the completion of medical records and the work pressure on the health workers involved, it was sometimes difficult to fill the gaps of any missing information, such as birth weights, retrospectively. Every effort was made by the management committee to minimise missing data, especially regarding maternal and neonatal outcomes.

Discussion

Summary

This initiative aimed to assess whether mothers were able to undertake fetal heart rate monitoring of their unborn babies immediately following the end of every contraction during labour in two rural public hospitals in Liberia, a country with extremely poor resources (both human and material). As 92% of mothers were able to undertake the monitoring themselves until their baby was born, the results indicate that this approach is feasible.

All of the comments and feedback from the participating women and adolescent girls (not just those selected in the Comment sections of the tables but also the complete set of comments provided as supplementary information), the overwhelming response (386 out of 400) was positive, with many comments reflecting how the monitoring helped a woman/
adolescent girl “feel strong”, cope with the pain of labour, or made her feel closer to her baby.

In our experience, it is most unusual for mothers in Liberia (as in many other low-resource settings where the hospital workforce is so limited and stretched) to be asked for their opinions on their experiences of labour and how it was managed. We asked for their opinions in an open way and did not raise the question as to whether they would prefer any alternative approaches to fetal monitoring, as stretched resources, lack of health workers, and subsequent high workloads, made such alternative approaches unfeasible.

Also, as explained in our Background, without this approach, many women would not have had her fetus routinely monitored because of the lack of, and high workload of, midwives.

To go through a pregnancy only to have a fetus or baby die because of a preventable condition that might have been detected by adequate fetal monitoring is a tragic and traumatic experience. This initiative may have helped to prevent some deaths while helping women/adolescent girls feel involved in what was happening to them and their babies during labour.

*Interpretation*

Since a study of mothers assisting midwives in monitoring the FHRs of their unborn babies has not been previously reported, as far as we can identify, it is difficult to compare our results with other studies.

This initiative confirmed that midwives responded to alerts from mothers regarding changes in FHR in a timely fashion and that trained, senior, health professionals (obstetric clinicians and doctors) were able to intervene appropriately and promptly. In 26 out of 461 women undertaking monitoring (in one undertaken with consent by a student midwife and the other by the midwife who took over from the mother who became tired), confirmed
changes in fetal heart rates were identified. Actions to improve the placental circulation (such as lateral tilt, oxygen and intravenous fluids), and, where possible, expedited delivery by vacuum or Caesarean section were undertaken following standard obstetric management as recommended by WHO [21].

All 26 neonates with changes in FHR survived, including 13 requiring resuscitation at birth. Ten neonates were admitted to the neonatal unit. One neonate developed birth asphyxia/HIE, but the immediate clinical outcomes of the 25 other neonates were good and all 26 were discharged home apparently well. However, long-term infant follow-up was not undertaken and should be in the future. The presence of neonatal clinicians and a functioning neonatal unit at CB Dunbar hospital were particularly valuable. Neonatal units that can provide advanced neonatal care by appropriately trained nursing staff (also a form of task-sharing) are now being established at 3 other rural county hospitals in Liberia.

In two mothers, the identification of FHR changes revealed previously unrecognized life-threatening obstetric complications. More experience may identify whether maternal FHR monitoring can consistently identify obstetric complications. Our results support Hofmeyr and colleagues [16] who stressed the importance of task-sharing in the integration of obstetric and neonatal care. Rapid access to effective obstetric management if FHR changes are identified must be ensured. Fetal monitoring in isolation has little value without an available health support system so that appropriate clinical intervention can be promptly undertaken. This situation may not be the case in some resource-limited settings, limiting the applicability of maternal fetal monitoring.

Limitations

Comparing figures with those from Tanzania [14] and from Sub-Saharan African countries [1, 12, 13,15], the absence of intrapartum stillbirths in the 461 fetuses monitored is
encouraging, as is the low prevalence of birth asphyxia/HIE. However, given no comparator, we cannot conclude that stillbirths and birth asphyxia/HIE can be prevented or reduced by maternal FHR monitoring. However, in reality, the lack of comparator data is represented by almost no fetal monitoring in public health hospitals in Liberia. We considered having a control group for this study but decided that given the situation described in the “Background”, such an approach would have been difficult. However, partway through this initiative, data on neonatal deaths and intrapartum stillbirths began to be collected by the Ministry of Health in Liberia. In the future, such data may provide control information for those maternity facilities where maternal FHR monitoring has not yet been undertaken.

Since we do not have robust baseline (comparator) data we cannot at this stage comment on whether Caesarean Section or operative vaginal delivery rates were increased or decreased by maternal FHR monitoring.

This initiative was conducted in a “real world” setting and suggests that maternal fetal monitoring can be incorporated into the daily work of a busy hospital maternity unit. The constant presence of obstetric clinicians was of considerable assistance, but such experienced health professionals may not always be available and alternative plans to address the health-workforce-shortage may be needed. Even with this additional cadre, communication and adherence to the study protocol were sometimes difficult, depending on the enthusiasm and involvement of all health professionals. Feedback of results to the midwifery workforce appeared helpful. Although we did not explicitly seek the views of attending midwives, which was a major limitation, the finding that midwives sometimes took over the monitoring if a woman was too tired or in too much pain to do it herself, shows their engagement in the process. More information from midwives could also be useful in helping to improve this technique, make it more sustainable and give a further
guide to the additional time needed to undertake maternal-fetal monitoring.

The potential for blaming the labouring women if there is an adverse outcome for their baby due to perceived failings in their self-monitoring is a very important consideration. However, there was no evidence that this situation happened, and, in our view, potential blame is unlikely as sadly, neonatal death and stillbirths are such common experiences in Liberia, that such events are almost expected.

We appreciate the frequency of monitoring can be onerous for some women, but there was no provision for birth partners or other relatives to be with women during labour, because of shortage of space and lack of privacy. We are planning to try and address this situation in the future.

**Developments and possible ways forward**

The long-term solution to fetal monitoring and improving the care of women in labour and their babies is undoubtedly to increase the availability of trained midwives. However, as explained in the “Background” to this paper, and especially the Global WHO data concerning midwives and nurses [2], given the current situation in Liberia and many other low income settings with minimal midwives and nurses a vast amount of resources and an increase is suitably trained health workers would be necessary to meet this goal, which is not possible in the short-term. However, once and if this situation is achieved, given the positive responses of participating mothers, we would continue to consider that the involvement of mothers in fetal monitoring may work well in collaboration with the enhanced role of skilled birth attendants.

The need for resuscitation in 8 of 461 (1.7%) of babies born without FHR changes identified by maternal monitoring, requires future research to ascertain whether changes are being missed or whether in some cases there are no measurable FHR changes identifiable by this monitoring technique in fetuses who subsequently require
resuscitation. In particular, it is important to know whether mothers and/or midwives can monitor reliably after every contraction with the same quality during the end of the first and second stages of labour when contractions can be much more painful. When appropriately undertaken and documented, with timely responses by care givers, the partograph is of major value in monitoring the progress of labour and safety of mother and her fetus. In support of Hofmeyr e.al.[16] who stressed the importance of task-sharing (in this case involving mothers), our findings show that it is feasible for pregnant women to undertake fetal heart monitoring during labour so that changes in FHR in relation to the end of every contraction are monitored. A decrease in FHR that persists after the end of a contraction, or continues after a contraction, is more likely to be pathological (Type 2 decelerations). WHO recommendations in place at the beginning this initiative [20,21] specified FHR documentation on the partograph every 30 minutes in the first stage and every 5 minutes in the second stage of labour and not with every uterine contraction, as in our initiative. Moreover, in these earlier guidelines [21, 22] WHO had recommended that the FHR be listened to immediately following the end of a contraction ONLY during 1) the initial assessment of the mother during labour; 2) when malpresentation or malposition are present; and 3) when inducing or augmenting labour. However, during the implementation of this initiative, in February 2018, WHO updated its guidelines on intermittent FHR auscultation in labour, now recommending auscultation every 15-30 minutes during the first stage, and every 5 minutes during the second stage of labour stating that auscultation should begin during a contraction and continue for at least 30 seconds after the contraction has ended.[7] These latest guidelines also recommend that if the FHR is not always within the normal range (110-160 bpm), auscultation should be prolonged to cover at least 3 contractions and also recommend recording the baseline FHR and the presence or absence of accelerations and
decelerations [7]. These updates in guidelines are welcome but, in our opinion, are quite complex, difficult to implement in facilities where there are few midwives, and do not concentrate on the most important time for intermittent auscultation, that is, at the end of every contraction. Our approach which involved auscultation from the end of every contraction, was easy to teach, convenient and feasible for individual mothers to undertake, and covered the most critical time for monitoring. For low resource settings, this approach may merit consideration by WHO and the wider international community. Future expansion of the project to include the possible role of female relatives and traditional birth attendants in supporting the mothers during monitoring; especially at times when contractions are particularly frequent and painful around the end of the first stage and during the second stages of labour, could be helpful. Unfortunately, the lack of space, privacy and resources such as toilets in the labour and delivery wards at present does not readily support this approach. Helpers such as traditional birth attendants or nurse aides may also be of value in collecting baseline/comparative data, and feedback from midwives.

An introduction to self-fetal monitoring during labour at antenatal visits would be beneficial, noting that a large proportion of pregnant women in rural Liberia do not always attend antenatal care.

The mothers’ comments on the lack of any pain control were of great concern and the mothers were clear about the need for pain control during labour. The absence of intrapartum analgesia made self-monitoring difficult for some women, which is partly the reason for listening to the fetal heart tones immediately after each contraction. However, a bradycardia that is slow to recover after a contraction may indicate fetal distress and this situation will be picked up by this form of monitoring.

We have now followed up the need for adequate pain control during labour within the
Liberian Ministry of Health and Social Welfare and have started a clinical audit on the use of intravenous paracetamol at CB Dunbar Hospital.

Some mothers stated that they would encourage relatives and friends to attend the hospitals involved in this initiative so that they might benefit from the fetal monitoring process. Given the encouraging findings of this study, we suggest that other hospitals in low resource settings, where there are few midwives and consequently, inconsistent fetal monitoring on the partograph, consider introducing maternal FHR monitoring, assisted by relatives where circumstances allow. Although a small minority of mothers expressed negative comments, there does not appear, to the best of our knowledge, to have been have any negative clinical consequences. Given the circumstances in countries where there are so few nurses and midwives, this approach supports realizing the rights of adolescent girls and women by enabling them to become more involved in their health care and the welfare of themselves and their unborn babies.

This initiative is continuing in both CH Rennie and CB Dunbar hospitals with further expansion planned in the near future into 3 additional rural maternity hospitals in Liberia.

Conclusions

The most promising findings of this initiative were the ability of the mothers to detect fetal heart rate changes and for midwives, obstetric clinicians and doctors to urgently respond to confirmed changes with appropriate clinical actions. The absence of intrapartum stillbirths, and the low rates of resuscitation and subsequent birth asphyxia/HIE in neonates who had been monitored are encouraging. A welcome finding was the positive comments made by 387 of 400 mothers about their involvement in the monitoring of their own unborn babies. If further studies in other settings, involving much larger numbers, continue to show these benefits, this new approach could help reduce the devastating problems of intrapartum stillbirth and birth asphyxia / HIE that are so
prevalent in low resource settings.

List Of Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| CBD          | CB Dunbar Hospital |
| CHR          | CH Rennie Hospital |
| CPAP         | Continuous Positive Airways Pressure |
| CS           | Caesarean Section |
| D50%         | Dextrose 50% intravenous solution |
| FHR          | Fetal Heart Rate |
| HIE          | Hypoxic Ischaemic Encephalopathy |
| IUFD         | Intra Uterine Fetal Death |
| IV           | Intra-Venous |
| LBNM         | Liberian Board for Nursing and Midwifery |
| MCAI         | Maternal and Childhealth Advocacy International |
| MOHSW        | Ministry of Health and Social Welfare |
| MW           | Midwife |
| NA           | Not appropriate or not available |
| NNU          | Neonatal Unit |
| NREB         | National Research Ethics Board |
| NS           | 0.9% saline intravenous solution |
| OC           | Obstetric Clinician |
| OR           | Operating Room |
| R/L          | Ringer Lactate intravenous solution |
| UNFPA        | United Nations Population Fund |
| UNICEF       | United Nations International Children's Emergency Fund |
| WHO          | World Health Organization |
Declarations

Ethics approval and consent to participate

The fetal monitoring project started as a service delivery intervention; when we realized the significance of the outcomes, we decided to approach it from a research angle for publication. As a programmatic intervention, it was fully within the domain of the Ministry of Health to implement. Re-framed as a research project, it was relevant to seek National Research Ethics Board approval.

In accordance with 45 CFR 46 the research was approved by the National Research Ethics Board (NREB) of Liberia through a full board review. The federal wide assurance number of the NREB is 000 21658 and organisation number 000 8374.

This approval was given after reviewing the documents relating to the initiative FHR monitoring project, which included the protocol stipulation that all mothers in labour (themselves individually and whatever their ages) would give their informed consent to be involved in the initiative (FHR monitoring project). Nine participants were aged under 16 years; despite their age, these young participants were deemed to be competent to give informed consent to be involved in the monitoring project as per the approved study protocol. Further consent from their parents or guardians was not sought. Because of poverty and complex social circumstances, it is common for mothers in Liberia to be aged under 16 years. Also, young mothers frequently attend the hospitals in labour by themselves, with no legally assigned family members in attendance and sometimes only a traditional birth attendant for support. Many women and adolescent girls do not know their age, and the illiteracy rate in Liberia is high (63%).

Consent for publication

Consent for Figure 1 was obtained from the patient using the MCAI consent form which is available on request.
Advice from a practicing lawyer revealed that there was no law in Liberia requiring explicit authorization for use of an anonymized quotation in any publication. Moreover, the consent, fetal monitoring, and birth summary forms, which included the comments made by mothers, was provided to the NREB as part of the application for approval allowing them to consider whether it was comprehensive/appropriate in making its decision to approve the study.

**Availability of data and materials**

The datasets used and/or analysed during the current study are available as two tables (one for CH Rennie Hospital and 1 for CB Dunbar Hospital) as 2 Additional Files.

**Competing interests**

The authors declare that they have no competing interests.

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None of the authors were paid to be involved with this present research project.

**Authors’ contributions**

The corresponding author affirms that all authors meet authorship criteria and that no others who have contributed have been excluded.

Trainee obstetric clinicians KB and NJ were responsible for overseeing consent for and collection of data at CB Dunbar and CH Rennie Hospitals respectively. They checked the drafts of the manuscript as they were prepared. They checked and approved the final version of the paper.

DS is the corresponding author and guarantor for the work. He is the voluntary medical
director of MCAI. He was jointly responsible for the concept, the design, and the implementation of the study. He was a member of the study management team. He was responsible for obtaining ethical approval for the study in collaboration with the Ministry of Health. He had overall responsibility for analysing the data. He wrote the original draft of the paper and was the lead editor for all subsequent drafts.

RM is the voluntary executive director of MCAI and Chair of Trustees and was jointly responsible with Professor Southall for the concept, the design, and the implementation of the study. She was a member of the study management team. She contributed to the analysis of the data, contributed to writing the original draft, and editing subsequent versions. She checked and approved the final version of the paper.

AAK was responsible for ensuring that the neonatal care (including neonatal resuscitation) was undertaken appropriately at CB Dunbar Hospital. He has recently helped MCAI establish a county neonatal unit at CH Rennie Hospital. He was responsible for the verification of all the data, especially in cases where FHR changes were detected and managed. He was a member of the study management team. He checked and approved the final version of the paper.

OD was the Medical Director at CB Dunbar Hospital during the first year of the project and was responsible for ensuring that the project was undertaken safely in this hospital. He provided clinical input into the writing of the paper and checked and approved the final version of the paper.

AM is the Medical Director at CH Rennie Hospital and responsible for ensuring that the project was undertaken safely in this hospital. He checked and approved the final version of the paper.

DW was responsible for researching the background to this research and is also the Special Advisor in Medical Ethics and Professional standards for MCAI. She is also a
volunteer international instructor of the Obstetric Clinician Training Programme in Liberia. She provided clinical input into the writing of the paper and checked and approved the final version of the paper.

MC is a volunteer international instructor of the Obstetric Clinician Training Programme in Liberia and works intermittently in CH Rennie Hospital. She was one of the main persons responsible for formulating the idea to undertake this project based on the high rates of stillbirth and early neonatal deaths in CH Rennie Hospital. She provided clinical input into the writing of the paper and checked and approved the final version of the paper.

BD was the Minister of Health for Liberia during the first year of the research project. She helped to ensure that the project was supported in the two hospitals. She helped with the design of the project and as Vice President of the College of Health Sciences, University of Liberia, provided comments on the draft of the paper and checked and approved the final version of the paper.

WJ has been the Minister of Health in Liberia since the beginning of 2018 and has continued to support the research project in the two hospitals. She also assisted with ensuring that the two hospitals were able to continue with the project following the change in Government of Liberia in January 2018 and has continued to support the further implementation of the project. She provided comments on the draft of the paper and checked and approved the final version of the paper.

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Tables

Table 1 Lowest and highest numbers of nurses and midwives per 10,000 of the population in 194 countries in the world. [2]

| 17 Countries with lowest numbers of midwives and nurses | Last Date of Data recorded | No. midwives and nurses per 10,000 population | 15 Countries with highest numbers of midwives and nurses | Last Date of Data Recorded | No. midwives and nurses per 10,000 population |
|--------------------------------------------------------|-----------------------------|---------------------------------------------|-------------------------------------------------------|-----------------------------|---------------------------------------------|
| Somalia                                                | 2014                        | 0.611                                       | Norway                                                | 2017                        | 181.247                                     |
| Liberia                                                | 2015                        | 1.007                                       | Switzerland                                           | 2016                        | 172.828                                     |
| Madagascar                                             | 2014                        | 1.039                                       | Iceland                                               | 2017                        | 156.806                                     |
| Central African Republic                               | 2015                        | 2.039                                       | Finland                                               | 2016                        | 147.230                                     |
| Malawi                                                 | 2016                        | 2.528                                       | Republic of Ireland                                   | 2016                        | 142.949                                     |
| Togo                                                   | 2015                        | 2.980                                       | Germany                                               | 2016                        | 131.967                                     |
| Bangladesh                                             | 2017                        | 3.067                                       | Luxembourg                                             | 2017                        | 123.496                                     |
| Dominican Republic                                     | 2017                        | 3.099                                       | Australia                                             | 2016                        | 126.612                                     |
| Niger                                                   | 2014                        | 3.109                                       | Uzbekistan                                            | 2014                        | 120.739                                     |
| Senegal                                                | 2016                        | 3.129                                       | Sweden                                                | 2016                        | 115.434                                     |
| Afghanistan                                            | 2014                        | 3.200                                       | Japan                                                 | 2016                        | 115.184                                     |
| Chad                                                   | 2016                        | 3.637                                       | Belarus                                               | 2014                        | 114.383                                     |
| Mali                                                   | 2016                        | 3.820                                       | Belgium                                               | 2016                        | 111.011                                     |
| Guinea                                                 | 2016                        | 3.844                                       | New Zealand                                           | 2017                        | 109.550                                     |
| United Republic of Tanzania                            | 2014                        | 4.126                                       | Denmark                                               | 2016                        | 103.004                                     |
| Mozambique                                             | 2017                        | 4.436                                       |                                                       |                             |                                             |
| Democratic Republic Congo (DRC)                        | 2013                        | 4.700                                       |                                                       |                             |                                             |

| Democratic Republic Congo (DRC)                        | 2013                        | 4.700                                       |                                                       |                             |                                             |

Monaco (Pop.38,000) and Niue (Pop. 1,600 excluded having high proportions but very small populations

Table 2 Questions to be addressed by this initiative.
| Questions to be addressed | Outcomes |
|--------------------------|----------|
| 1. Can women in labour be educated to detect changes in FHR (especially fetal bradycardia and fetal tachycardia) through self-monitoring of the FHR with a doppler monitor and alert the attending midwife? | 1a) Number of women (out of all approached for consent) hospital maternity units willing to participate in the self-monitoring initiative 1b) Number of women (out of all approached for consent) in the self-monitoring initiative who were able to detect which were confirmed as abnormal and possibly harmful by the clinically attending midwife. |
| 1. Do attending midwives respond to alerts from women in labour regarding suspected FHR changes and do they initiate the agreed course of action (the birth asphyxia and stillbirth prevention protocol) every time in a timely manner? | 2a) Number of times (out of all possible) an attending midwife responded to an alert from a participating woman in labour detecting potentially harmful change in FHR. 2b) Action taken by an attending midwife responding to an alert from a woman in labour of a possible FHR change and whether the agreed birth asphyxia and stillbirth prevention protocol had been followed. |
| 1. Did the labouring women find the experience of monitoring their unborn babies helpful? | 3a) How many mothers found the monitoring helpful 3b) How many mothers found the monitoring difficult 3c) How many mothers had to discontinue the monitoring |
| 1. What measure could be implemented to improve the attainments of the above first three objectives and result in a sustainable programme? | 4a) Improvements in obtaining consent 4b) Improvements in the documentation of changes 4c) Feedback of results to the midwives on the maternity wards 4d) How to achieve sustainability given the temporary availability of trainee obstetric clinicians |
| 1. Was the attending midwife able to initiate an immediate course of treatment when she was alerted by a woman in labour who had identified changes in her FHR? 2. Were professionals trained in advanced obstetrics and neonatal care able to provide effective treatment when asked to by the attending midwife? | Treatments given and outcomes given in the mother Treatments given and outcomes in the newborn infant |

Table 3 Selected maternal comments from participants at CB Dunbar and CH Rennie Hospitals
| Case Number | Comment |
|-------------|---------|
| 1 CHR       | The monitoring was fine, it gave me courage to go through my pain knowing my baby was fine. |
| 34 CHR      | She further stated due to the exercise she will always come to CH Rennie for maternity care during pregnancy. |
| 35 CHR      | I felt that I am important when you told me to be a part of my baby monitoring process. It helps me a lot. |
| 37 CHR      | The monitoring was good. It help even us that cannot read or write listen to our own baby. |
| 40 CHR      | I thank god for the programme I am happy to hear my baby heart-beat. Please continue it. |
| 43 CHR      | I am comfortable doing this as it helped me form part of my baby monitoring process. |
| 61 CHR      | Listening to my baby heart sound was very helpful to me. I felt that my right was respected as I took in the monitoring. Thanks for this program. I am happy. |
| 62 CHR      | I am happy to hear my baby heart. I knew that I was carrying a live baby in my womb. |
| 70 CHR      | Thank you for this. It help me but I was in pain and so it make me angry first but I overcome it later (age 35). |
| 95 CHR      | Getting involved in the process is something amazing to me. I felt part of my care and thank God that I am important. |
| 156 CHR     | I feel important in the coming of my baby, This modern method is very important it help a lot thank you. |
| 158 CHR     | I like it so much doctor that real good thing the government put in place here. I will tell all my sisters that come to the hospital. |
| 4 CBD       | According to mum it is a good step to do because it helps you to notice danger sooner. |
| 51 CBD      | I found the monitoring helpful it helps me go through my pain. |
| 52 CBD      | I felt good listening to my baby it helped me to learn a new thing. |
| 123 CBD     | According to mum, this is the first time seeing patient to be working for herself. She said it is a good thing in labour is bad because of the pain. |
| 133 CBD     | Patient said she’s very happy because she seen baby breathing well and she herself okay. According to her she pregnant she will come and give birth to CB Dunbar hospital. |
| 138 CBD     | According to mum, she love the procedure but is not easy to go through. |
| 139 CBD     | According to mum, she love the idea because other pregnant women goes to the hospital and comes back with their hands it looks sorry full. |
| 151 CBD     | Mother said she found the monitoring helpful in that she has a live baby. She was cooperative and was a mother to join the process. |
| 162 CBD     | Patient admitted that was good thing for herself to listen to her baby heart-beat. It made her believe that breathe inside her mother’s womb. |
| 169 CBD     | Mother was happy to hear her baby heart beat because she stay in labour for long and worry about her baby wellbeing. |
| 180 CBD     | According to patient she was surprised to know that baby heart can beat in the mother stomach and it helps about her baby wellbeing. |
| 200 CBD     | I like to have the same chance to listen to my unborn baby the next time I am in labour. |
| 203 CBD     | I enjoy listening to my baby but my next labour there should be pain medicine for labour. |
| 239 CBD     | It help me to put more effort for my baby. To know that my baby is still living in my stomach. |
| 242 CBD     | It is hard to be in pain and monitor your baby. You must be doing it for us. Thank God my baby is living! The machine can cause more pain on the stomach. |
| 255 CBD     | It help me because it did not allow me to go to surgery. It help me because my baby was born alive and vaginal delivery. It help me so much even though is more difficult to do but I try doing to have got good breech delivery. |
| 274 CBD     | It is very good and helpful to me. At least all “big bellies” should know how to do the monitoring before it hurt. |
| 275 CBD     | I like the monitoring it make my baby live. No problem with the monitoring. It only hard to hold the machine without stomach hurting. |

Legend to Table 3. Additional maternal comments following changes in fetal heart rate are reported in Tables 4 and 6 below and in the data repository in the Additional Files. Abbreviations are defined in the list given earlier in the manuscript.
Table 4 Clinical information and outcomes of FHR changes identified by maternal monitoring. Abbreviations: see list

| Hospital and number | Maternal age (years) and parity | Change in FHR identified | Action taken | Apgar scores at 1 and 5 minutes | Resuscitation given |
|---------------------|--------------------------------|--------------------------|--------------|-------------------------------|---------------------|
| CHR 46              | 29, G5P2                        | By mother. FHR 115 plus meconium Confirmed by MW | Lateral tilt and intravenous cannula with NS bolus Vacuum delivery | 9 and 10 | None |
| CHR 50              | 34, G3P2                        | By MW and mother during training in the use of the sonicaid at time of admission.. No FHR was identified and there was meconium | Ultrasound confirmed IUFD. Vacuum delivery was undertaken | NA | NA |
| CHR 99              | 17, G2P0                        | By mother at 46th contraction FHR 109 with meconium | Cervix fully dilated and urged to push | 4 and 7 | Yes. Bag and mask ventilation, adrenaline and chest compressions for 10 minutes. Admitted to the NNU for post resus care and close monitoring Developed convulsions due to HIE and treated successfully with phenobarbital and recovered and was feeding normally at discharge home aged 7 days. |
| CHR 102             | 19, G2P0                        | By mother FHR 119 at 49th contraction. There was + meconium present | Vacuum delivery | 7 and 10 | No |
| CHR 133             | 23, G2P1                        | By mother FHR 119, 117, 116. No meconium. Patient was not progressing at this stage. 2 cm cervical dilatation with mild contractions. | MW/OC took over the monitoring due to the bradycardia. Doctor contacted. Patient was laterally tilted, given oxygen, D50%, hydrated and rushed to the OR for CS. | 7 and 10 | No |
| CHR 135             | 24, G2P1 No previous CS         | By mother FHR 163-165 with meconium. Signs of Bandl’s ring and obstructed labour with haematuria identified. | Not receiving oxytocin. Emergency CS | 9 and 10 | No |
| CHR 136             | 26, G3P2                        | By mother FHR 119,110,118. No meconium. OC and doctor contacted and confirmed bradycardia | Given facial oxygen, lateral tilt, N/S and D50%. Patient was 6cm dilated at this | 8 and 10 | No |
| CBD    | G1P0 | G2P1 | G3P1 | G3P2 |
|--------|------|------|------|------|
| CHR 157 | 19   | 17   | 22   | 28   |
| stage. Emergency CS | stage. Emergency CS | stage. Emergency CS | stage. Emergency CS |
| By mother at 46th contraction FHR 117, then 114, then 116, then 113. No meconium. Fully dilated but descent only minus 2 | FHR found to be 95-100 by mother, FHR was repeated by midwife and confirmed low, 95-98, and Doctor on call was also informed. | Mother reported a change in FHR but when checked by MW found FHR to be normal at 142. Meconium was present | Mother noted change and contacted MW on 15th contraction. MW noted FHR 118 and informed OC. Meconium was present repeat fetal heart rate was 105. |
| Lateral tilt, D50%, oxygen, NS and FHR still below 120 and when head reached 0 station re: ischial spines after 10 minutes and then vacuum delivery | Patient was placed in a left lateral tilt position Patient was reviewed and decision to CS was taken for fetal distress plus prolonged labour | Doctor informed but no action was considered necessary | Mother put in lateral tilt position and informed Dr who reviewed patient and found fetal heart rates 110, 105, and 108. Emergency CS was performed |
| 7 and 10 | 6 and 9 | 6 and 10 | 8 and 10 |
| No | None | None | None |
| 5 and 7 | 6 and 9 | 6 and 10 | |
| Bag and mask ventilation. Admitted NNU for 5 days and treated for sepsis. | Bag and mask ventilation and admitted to NNU. No HIE and went home. | This baby was resuscitated for 5 minutes with bag and mask ventilation and then transferred to the NNU where he was immediately placed on nasal CPAP and an IV line was opened to serve | |
| CBD 128 | 16, G1P0 | On the 14th contraction the mother called the MW because the FHR was low. The MW confirmed FHR 98, called for help and undertook lateral tilt. Meconium was present. | The OC was contacted. She opened IV line and gave R/L 1000 mL, informed the doctor on call. The doctor came and assessed the patient and said we should prepare patient for CS. CS was done for prolonged labour and abnormal FHR. | 5 and 10 | Neonate was resuscitated for 7 minutes by bag and mask ventilation before transferring to the NN. She was placed on nasal CPAP for 24 hrs and was also managed for risk of sepsis. Neonate improved after 8 days and was discharged. |
| CBD 131 | 15, G1P0 | On the 7th contraction, mother detected fetal bradycardia. MW called and checked and confirmed FHR 105. Meconium was present. Grade 3 OC was called. | Lateral tilt was undertaken and fast vaginal delivery arranged as 9cm cervix dilated. Birth weight 1.9Kg small for dates. | 7 and 10 | Baby was resuscitated for 2 minutes by bag and mask ventilation and then transferred to NN. She was placed on nasal CPAP for 24hrs and patient condition improved. Baby was also managed for risk of neonatal sepsis because mother’s amniotic fluid was purulent, foul-smelling during delivery. The baby was discharged home after 10 days with a weight of 2.3kg |
| CBD 147 | 28, G5P4 | On 6th contraction, mother detected bradycardia. MW confirmed FHR 108. Meconium was present. | OC contacted. Lateral tilt performed. IV cannula inserted and given NS 500ml. Normal vaginal delivery occurred. | 5 and 8 Male | Bag and mask ventilation given. No HIE occurred but he needed 5 days of antibiotics for umbilical infection. |
| CBD 153 | 32, G5P3 | On 2nd contraction monitored, Mother identified rapid heart rate. MW confirmed FHR 190 and called for help. | Doctor called and attended. Lateral tilt and IV cannula and N/S 500ml set up. Vacuum delivery was undertaken. | 6 and 8 | Neonatal clinician was called and baby resuscitated with bag and mask ventilation and recovered within 1 minute. Responded well and taken to |
| CBD 158 | 17, G2P1 | On 6th contraction, Mother reported fall in HR. MW confirmed FHR 109. Meconium present. | Lateral tilt applied and IV cannula inserted with R/L 500mls plus Dextrose 50% 30ml. OC contacted and quickly delivered the baby vaginally. | 6 and 7 | Mildly depressed but no resuscitation needed. Neonatal clinician continued monitoring and care. |
| CBD 160 | 26, G3P0 | On 27th contraction, Mother detected slowing of FHR. MW confirmed FHR 109. Grade 2 meconium was present. Dr on call contacted. | Lateral tilt and IV cannula inserted. R/L 500mls given IV. Doctor arrived and undertook CS. | 7 and 10 | Resuscitated for 2 minutes with bag and mask ventilation. |
| CBD 169 | 16, G1P0 | On 7th contraction mother noted fast heart rate. MW confirmed FHR 167. Patient came in fully dilated but evidence of obstructed labour due to persistent occipito-posterior malposition. | Lateral tilt and IV cannula inserted. NS 500mls given IV. Doctor arrived and undertook CS. | 9 and 10 | None needed |
| CBD 172 | 42, G9P8 | On the 7th contraction mother noted a slow heart rate. MW confirmed FHR 102. Meconium was present and a cord prolapse identified. | The OC was notified and implemented knee chest position and inserted NS 300mls into the bladder to reduce cord compression. IV cannula was inserted and NS 500mls given. A CS was then undertaken. | 6 and 10 | Depressed breathing. Resuscitated for 1-3 mins with bag and mask ventilation. Taken to NNU as 30 weeks' gestation No HIE. Home after 14 days |
| CBD 177 | 17, G2P1 previous CS | On 12th contraction, MW reported a FHR 124. Meconium present. FHR then dropped to 119. | OC was called and after lateral tilt established IV line and gave 500ml NS. A CS was then undertaken. | 7 and 8 | No resuscitation needed but foul-smelling amniotic fluid at CS led to NNU admission and IV antibiotics. |
| CBD 188 | 32, G1P0 | On the 8th contraction mother noted a slow heart rate. MW confirmed FHR 110. Meconium was present. OC informed and FHR was 112. Cervix fully dilated. | Lateral tilt and placed in delivery room for vacuum delivery. However, within 5 minutes delivered spontaneously. A very short umbilical cord was present. | 5 and 7 | Depressed breathing Resuscitated for 5 mins with bag and mask ventilation and taken to NNU and given antibiotics. Later became stable and discharged. |
| CBD 235 | 31, G5P4 | On the 30th contraction mother noted a slow heart rate. MW confirmed FHR 118. | MW performed lateral tilt and informed the OC and set up IV infusion of R/L 500ml. Dr ordered repeat and FHR 106. Cervix only 4cm dilated. Descent 3 / 5. Discussion for CS was done but no CS materials available so patient was referred to another hospital. | 8 and 9 | None needed after CS at referral hospital |
| CBD 251 | 19, G1P0 | On the 20th contraction OC and student MW noted a slow FHR 105. No meconium seen. | Lateral tilt was undertaken. The cervix was already 10 cm dilated and there were poor maternal efforts. An IV cannula was inserted and she was given 30 ml dextrose 50%. Baby was delivered by vacuum. | 5 and 6 | Yes by neonatal clinician bag and mask ventilation for 5-10 mins. Admitted to NNU for neonatal depression. Neonate recovered quickly on nasal CPAP. Improved and went home well. |
| CBD 272 | 19, G1P0 | On 30th contraction mother noted slowing of FHR. There was no meconium at this time. MW and OC identified FHR of 115, 118,122. | Lateral tilt and Doctor notified. An IV cannula inserted and given N saline 500ml plus Dextrose 50% 30ml. The cervix was 10cm dilated. OC did vacuum with Dr present but failed 3 times. Dr and OC proceeded to immediate CS. Intraoperative meconium was present | 5 and 7 | Bag and mask ventilation for mild respiratory depression. Recovered rapidly and went home. |
| CBD 273 | 22, G4P0 | On 51st contraction mother noted slowing of fetal heart rates. MW recorded FHR 109, 178,120,110,181,102,130. Meconium was present | Lateral tilt was performed, and OC notified. IV fluids were started and 30 ml of 50% dextrose given IV. The doctor was also called and due to FHR changes, high station O, and bad obstetric history (G4P0) proceeded with the OC to CS. | 8 and 10 | None |

Abbreviations are defined in the list given earlier in the manuscript.

Table 5  Clinical information and outcomes in newborn infants needing resuscitation at birth where mothers had declined consent to participate in the FHR monitoring. For abbreviations see list.
| Hospital and number | Maternal age (years) and parity | Change in FHR identified on partograph | Delivery | Apgar scores at 1 and 5 minutes; Wt. of baby | Resuscitation given | Maternal comment | Other inform: |
|---------------------|---------------------------------|---------------------------------------|----------|---------------------------------------------|--------------------|-----------------|---------------|
| CBD 71              | 32, G3P2                        | None                                  | Normal vaginal delivery | 2 and 3 3.4Kg | Resuscitated with bag and mask ventilation, chest compressions and oxygen | Admitted NNU but later died aged 2 days from HIE | None |
| CBD 184             | 25, G2P1                        | None                                  | Normal vaginal delivery | 4 and 6 Depressed 2.8Kg | Resuscitation was done with bag and mask ventilation and was taken to the neonatal ward. Treated with antibiotics. Outcome was good and discharged. | None | Patient contin rate m though the first |
| CBD 192             | 18, G1P0                        | None                                  | Normal vaginal delivery | 2 and 6 Very depressed | Resuscitated by bag and mask ventilation by neonatal clinician | Died aged 3 days from HIE | None | Patient contin rate m though the first |

Abbreviations are defined in the list given earlier in the manuscript.

**Table 6** Clinical information and outcomes in newborn infants needing resuscitation at birth where monitoring had not identified any FHR changes
| Hospital number | Maternal age (years) and parity | Change in FHR identified | Delivery | Apgar scores at 1 and 5 minutes; Wt. of baby | Resuscitation given | Maternal comment |
|-----------------|-------------------------------|--------------------------|----------|---------------------------------------------|---------------------|-----------------|
| CBD 164         | 25, G2P1                      | None. Monitored only every 30 minutes immediately following 14 contractions | Preterm labour and normal vaginal delivery | 5 and 7 depressed at birth 1.8 Kg | Neonatal clinician called, resuscitated with bag and mask for 12 minutes and taken to neonatal ward | No HIE |
|                 |                               |                          |          |                                             | Mother said the monitoring help her with her baby. She is willing and cooperates with other mothers. |
| CBD 176         | 18, G1P0                      | None. Monitored only every 30 minutes immediately following 13 contractions | Normal vaginal delivery | 5 and 10 depressed at birth 3.9Kg | Neonatal clinician was called, did 10-15 mins bag and mask ventilation. Oxygen saturation 54%. Admitted NNU. No HIE and went home aged 7 days | According to mother monitoring is good deliver her baby to interested in doing it |
| CBD 179         | 18, G2P1                      | None. Monitored only every 30 minutes immediately following 11 contractions | Normal vaginal delivery | 7 and 10 2.8 Kg | Bag and mask ventilation used for 5 minutes and then recovered. No HIE | |
| CBD 224         | 29, G2P1                      | No abnormality detected following 12 contractions | Vacuum delivery unable to push | 5 and 8 3.9Kg | Resuscitated by bag and mask for 10 minutes. Admitted to NNU and given 7 days antibiotics. No HIE. | It help because with pain I refused to listen to them. I still got a healthy baby |
| CBD 238         | 23, G2P1                      | No abnormality detected following 15 contractions | Normal vaginal delivery | 2 and 0 1.3 Kg 34 weeks’ gestation | Resuscitated by bag and mask ventilation plus chest compressions for 25 minutes. But then died. | I like the monitoring during labour |
| CBD 240         | 15, G1P0                      | No abnormality detected following 58 contractions | Normal vaginal delivery | 5 and 7 2.5Kg | Resuscitated by bag and mask ventilation by neonatal clinician for 8 minutes then improved and discharged. No HIE. | I see the monitoring of my baby because it still living. Patient is not literate |
| CBD 243         | 23, Gravida G2P1              | No abnormality detected following 23 contractions | Vacuum for reduced maternal effort | 6 and 8 3.1 Kg | Bag and mask resuscitation for 7 minutes. Baby was admitted to the NNU for observation. No HIE but had malaria and was treated for 10 days and then discharged well. | I like the monitoring of me and baby I make to him, the baby is still living. Patient is not literate |
| CBD 286         | 23, Gravida G1P0              | No abnormality detected following 12 contractions | Vacuum for exhaustion: couldn't push | 7 and 8 3.3 Kg | Bag and mask resuscitation one-two breaths only before baby breathed. Not admitted to NNU. | I find it good. It helps because my baby problem with it. |

Abbreviations are defined in the list given earlier in the manuscript.

Additional Files

Additional File (PDF) Master copy of the consent form, fetal heart rate monitoring and data
collection and maternal comment form.

The consent form provides the information that was either read by each participating mother, if literate, or read to the mother by the obstetric clinician. The form could either be signed by the mother or she could provide her fingerprint from an ink pad.

The FHR monitoring form was given to each mother who was asked to tick each time she listened to the FHR for approximately one minute following the end of every uterine contraction. If the mother considered that the FHR had changed, she would notify the midwife caring for her who would check the FHR, count it, and write down her own findings on this chart.

The data collection form recorded a summary of potentially relevant clinical data on the labour, delivery, neonatal Apgar Scores at 1 and 5 minutes, and any resuscitation given to the baby. Recorded on this form was any admission of the neonate to the neonatal unit and any treatment given to the baby. The form also includes a section in which the comments made by mothers on their experience of the monitoring were recorded either by themselves or as transcribed by the attending midwife or obstetric clinician.

The Additional files also contain two tables (one for CH Rennie Hospital and 1 for CB Dunbar Hospital) summarizing the clinical data collected on all mothers approached for consent, together with the comments made on the monitoring process by the mother, either written directly or transcribed for illiterate mothers by the obstetric clinician completing the form and outcomes of the newborn babies.

Figures
Figure 1

Mother undertaking fetal monitoring
Figure 2

Time lines of major inputs to the initiative

| TIMINGS                  | 31ST JULY 2017 | 24TH OCTOBER 2017 | 19TH February 2018 | 21ST MARCH 2018 | 27TH APRIL 2018 | 8TH May 2018 | 24TH OCTOBER 2018 |
|--------------------------|----------------|-------------------|--------------------|-----------------|-----------------|---------------|--------------------|
| Project active and reported so far in the manuscript |                |                   |                    |                 |                 |               |                    |
| Project undertaken in CB Dunbar Hospital               |                |                   |                    |                 |                 |               |                    |
| Project undertaken in CH Rennie Hospital                |                |                   |                    |                 |                 |               |                    |
| Request for written or verbal comments from mothers     |                |                   |                    |                 |                 |               |                    |
| Application to NREB                                      |                |                   |                    |                 |                 |               |                    |
| Request for written rather than informed verbal consent from mothers to participate |                |                   |                    |                 |                 |               |                    |
| Successive improvements in data monitoring forms completed by mothers |                |                   |                    |                 |                 |               |                    |
| Communication problem where mothers at CB Dunbar monitored FHR every 30 minutes following contractions rather than after every contraction |                |                   |                    |                 |                 |               |                    |

Figure 3

Flow chart showing consent and comments for this initiative
Supplementary Files

This is a list of supplementary files associated with the primary manuscript. Click to download.

MASTER CB Dunbar Hospital Table anonymised for repository .docx
Revised SQUIRE2.0_Checklist completed.docx
Additional file for BMC paper.pdf
MASTER CH Rennie Table anonymised for repository.docx