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Sonographic Evaluation of the Mechanism of Active Labor (SonoLabor Study): observational study protocol regarding the implementation of the sonopartogram

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ABSTRACT

Introduction Over the last decades, a large body of literature has shown that intrapartum clinical digital pelvic estimations of fetal head position, station and progression in the pelvic canal are less accurate, compared with ultrasound (US) scan. Given the increasing evidence regarding the advantages of using US to evaluate the mechanism of labour, our study protocol aims to develop sonopartograms for fetal cephalic presentations. They will allow for a more objective evaluation of labour progression than the traditional labour monitoring, which could enable more rapid decisions regarding the mode of delivery.

Methods/analysis This is a prospective observational study performed in three university hospitals, with an unselected population of women admitted in labour at term. Both clinical and US evaluations will be performed assessing fetal head position, descent and rotation. Specific US parameters regarding fetal head position, station and progression in the pelvic canal will be recorded to develop nomograms in a similar way that partograms were developed. The primary outcome is to develop nomograms for the longitudinal US assessment of labour in unselected nulliparous and multiparous women with fetal cephalic presentation. The secondary aims are to assess the sonopartogram differences in occiput anterior and posterior deliveries, to compare the labour trend from our research with the classic and other recent partogram models and to investigate the capability of the US labour monitoring to predict the outcome of spontaneous vaginal delivery.

Ethics and dissemination All protocols and the informed consent form comply with the Ministry of Health and the professional society ethics guidelines. University ethics committees approved the study protocol. The trial results will be published in peer-reviewed journals and at the conference presentations. The study will be implemented and reported in line with the Strengthening the Reporting of Observational Studies in Epidemiology statement.

Trial registration number ClinicalTrials.gov Registry (NCT02326077).

Strengths and limitations of this study

- The multicentre design on representative population and the blinded clinical/ultrasound assessment aim to intercept the potential sources of bias.
- The SonoLabor Study differs from previous studies as it aims to assess all stages of labour, rather than just the second stage of labour to elaborate nomograms for the longitudinal ultrasound assessment of labour in unselected low-risk population, an important issue of the future sonopartograms.
- Sonographic and clinical evaluation of the labour progression in any cephalic presentation (not only with occiput anterior position).
- Our study aims to develop curves for labour monitoring which will be not only objective, but also adapted to contemporary practice. Clinical studies showed that the pattern of labour progression and the present characteristics of the partogram differ significantly from the traditional Friedman curve.
- The high number of labouring women needed to investigate the characteristics of each clinical situation targeted in the study design. In order to produce specific sonopartograms regarding the maternal characteristics and occiput position, an important number of patients would be required, that may not achievable during our present research. We hope that the publication of our study protocol, dissemination of the results and the storage of the anonymised collected data in a research depository will serve to future larger studies that will help to collect or complete the necessary data.
- The concept of normality is population based and depends on various management attitudes (for example, epidural analgesia, active management of labour), different characteristics of the partogram are observed that may affect generalisability.
INTRODUCTION

Over the last decades, a large body of literature has shown that clinical digital pelvic estimations of fetal head position, station and progression in the pelvic canal are not accurate during the first and second stage of labor, poorly reproducible when compared with ultrasound (US), poorly reliable, experience dependent and often inexact in challenging labour circumstances, such as: prolonged first stage of labour, cases with arrested cervical dilatation, obstructed labour, fetal head engagement, posterior and transverse occiput locations, or caput. This may imply significant consequences on the decision of the appropriate delivery mode, because digital examination is less reliable especially when obstetrical interventions are more likely to be needed. Intrapartum sonographic evaluation may not provide a solution for all these conditions mentioned above, but previous studies have demonstrated the potential to decrease the rate of late caesarean extractions in prolonged labour cases, and the various approaches proposed in the literature were considered by our study design. Many studies provided sonographic data regarding the fetal head descent/progression (FHPr) in the second stage of labour and proposed several easily measurable and reliable parameters, capable to predict the vaginal or operative outcome of the delivery with occiput anterior positions. The literature regarding US evaluation in the first stage of labour is less, but based on available data US evaluation appears to be useful for the prognosis of labour.

Given the increasing evidence regarding the advantages offered by using US in labour, our group concluded that the development of a sonopartogram, as an adjuvant to or a replacement of traditional labour monitoring, provides the setting for a more objective evaluation of labour progression, which would enable more rapid decisions regarding the mode of delivery. Intrapartum US evaluation is not meant to change the standard principles for labour mechanism evaluation, but to provide accurate evaluation of the main parameters involved: fetal head position and rotation, fetal head progression and engagement.

The SonoLabor Study aims to provide new objective evidence regarding the evaluation of the mechanism of labour with US. There is little information in the literature regarding the ultrasonographic monitoring of the entire active labour. A recent proof-of-concept study showed that the sonopartogram is feasible in most cases. However, a study of the paired clinical and sonographic assessments of labour in a large, unselected population has not yet been conducted. Furthermore, there are no nomograms for the US monitoring of labour. Nowadays, the use of US in labour is generally limited to research settings and a relatively small number of women have been studied. Therefore, efforts should be made to describe the value of an objective partogram in general practice.

This study is designed to produce an original multicentre longitudinal assessment of the mechanism of active labour, including both stages, in a representative population, using concomitant blinded clinical and sonographic evaluations in unselected low-risk parturient women at term. A unique point of our protocol is the comparative evaluation of the US parameters for various clinical situations. This may facilitate the use of different nomograms in labour, adapted to the clinical characteristics of the labouring woman. Another strength of our study is the multicentre design that is useful to achieve a proper study size and an opportunity to compare the data recorded in different settings.

The aim of this paper is to describe the protocol of the study.

Objectives

The primary objective of this study is the development of nomograms for US-measured variables during labour in unselected nulliparous and multiparous women at term with fetal cephalic presentation.

The secondary objectives of the study are the following:

- To compare the US pattern of labour evolution in nulliparous and multiparous women.
- To study the influence of occiput position, body mass index (BMI), parturient age on the labour progression evaluated by US.
- To correlate the labour trend from our study with the Friedman studies and other recent research on the partogram regarding the progression of labour by means of objective US evaluation.
- To correlate the US and standard clinical findings regarding the mechanism of labour, for example, fetal occiput position and head descent during active labour.
- To investigate the correlations between the data of the participating centres.
- To analyse the temporal variation of the sonographic measurements in spontaneous vaginal delivery versus obstructed labour in nulliparae versus multiparae.
- To analyse the evolution of the sonographic measurements in spontaneous vaginal delivery versus obstructed labour in fetuses with occiput anterior versus those with persistent occiput posterior.
- To investigate the value of combined US measurements to predict the outcome of vaginal delivery.

METHODS AND ANALYSIS

Study design and setting

This is an observational cohort prospective study, which will take place in three tertiary maternity hospitals (University Emergency County Hospital Graiova, Alexandra University Hospital of Athens and Ippokrateion Hospital Thessaloniki), with more than 4000 deliveries per annum. The study aims to record simultaneously the labour progress by clinical and US evaluations in women in labour at term, with singleton cephalic presentation. We will include low-risk pregnancies, according to the criteria defined in the Participants section.

Patient and public involvement

We conducted a previous study during the development of this research question, where we evaluated the acceptability of the method and found that the vast majority of
labouring women (98%) agree with the supplementary US investigation protocol and the demographic characteristics did not influence the rate of acceptance. Most of the women (93% of accepters and 75% of decliners) had little difficulty deciding whether or not to have the scan protocol. All women who were scanned during labour found it an acceptable experience, and only 21% of women without epidural anaesthesia rated the perceived difficulty as ‘mild’ or ‘discomforting’. Women rated having the intrapartum scan as being significantly less difficult than having a cervical smear, transvaginal scan or having a digital clinical evaluation. Two-thirds (67%) of the patients expressed increased confidence while being able to follow along the medical personnel the FHPr on the US screen. Almost all of the consenting women (97%) who had the intrapartum US scans and all the four decliners said they would definitely or probably agree such US monitoring in a future labour, if this technique is proven useful for the labour outcome.

Participants
All pregnant women admitted in active labour at term are considered eligible for the study. They will be consecutively included in the study, depending on the availability of the US operators involved in the study. We will try to attract a large team of collaborators, in order to investigate as many eligible cases as possible. Cases planned for elective caesarean section, or involving imminent delivery, with non-elective caesarean section, or involving imminent artistic rupture of the membranes, oxytocin augmentation in the first stage of labour, when instrumental delivery is attempted, as presented in the Interventions section. The only direct benefit of the labouring women who participate in the study would be the communication between the obstetrician and sonographer regarding the FHPr on the US screen. Almost all of the consenting women (97%) who had the intrapartum US scans and all the four decliners said they would definitely or probably agree such US monitoring in a future labour, if this technique is proven useful for the labour outcome.

Interventions
All pregnant women who meet the inclusion criteria will be assessed clinically by the physician on duty. The managing clinician is a senior consultant not involved in the study.

Clinical examinations will take place in women in active labour just before the US assessments (figure 1). The clinician will note the observations on a specially designed partogram-like sheet that will be used for women in labour who agree to participate in the study. The following labour parameters must be noted before US assessment:

- Cervical dilation in centimetres.
- FHPr—determined by the evaluation of head station in relation to the ischial spines.
- Presence of caput, with the approximate diameter.
- Presence of moulding and grading: closure of sutures measuring the midline angle.
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Clinical examination will be followed by transabdominal and transperineal US evaluations conducted by obstetricians with appropriate training in US in labour, with minimum 1 year of experience in the field. Mobile and compact US machines will be used: Logic e (GE Healthcare, China), GE Voluson P6, Samsung R7, BenQ T3300 and ALOKA f31 equipped with 2–5 and 2–6 MHz 2D convex transducers.

The objectives of US evaluations are similar to those of standard clinical assessment. The purpose of US evaluations is to document the progression of labour using objective measurements for the main parameters involved in the mechanism of labour:

- FHPr, by determining occiput position.
- FHPo, by determining occiput position.
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Procedures
Recruitment
During their usual consultation in the labour ward, in an eligible case, the physician on duty provides brief information about the research and invites the patient to take part in the study. If the patient shows interest in the study and meets the inclusion criteria, a face-to-face appointment with the US operator is arranged. The details of the study and the potential benefits of the research will be thoroughly explained to the patient. The only direct benefit of the labouring women who participate in the study would be the communication between the obstetrician and sonographer regarding the FHPr on the US screen. Almost all of the consenting women (97%) who had the intrapartum US scans and all the four decliners said they would definitely or probably agree such US monitoring in a future labour, if this technique is proven useful for the labour outcome.
In table 1, we present the sonographic measurements, in relation to the acquisition planes and the features of labour mechanism that are involved. The images will be stored on the hard disk drive of the system for offline analysis and the measurements will be performed by the sonographer who evaluated the case, according to the techniques described in the literature (table 1).

Because of ethical issues, the design of the study states that the attending obstetrician should be informed in case of clinical and US discordance when instrumental or operative delivery is attempted. The digital evaluation is considered to be correct if the FHPo is within ±45° of the US determination.

Cervical dilation will be evaluated only clinically, as the evaluation of this parameter is best achieved with digital assessment.31 Timing: clinical and US scans is performed hourly until complete dilation (first phase of active labour) and at every 15 min after complete dilation (second phase). The purpose of the apparently high number of examinations was to obtain accurate information in each labour in terms of correlation of FHPr, FHPo and cervical dilatation. In a previous study in our clinic, this methodology proved to be acceptable for the parturients.40

The frequent evaluations in the second stage are meant to offer a better analysis of this critical stage of labour. Notification of time delivery will be used to calculate the time interval from each scan to delivery.

US images will be saved and stored on US hard disk during labour, then transferred by the sonographer to a designated personal computer storage unit after birth. The images will be reviewed during the following week, by the same sonographer, who will also input the offline measurement results into the database.

The sonographer and the clinician are blinded to each other’s findings (except FHPo, during the circumstances mentioned above) as the specific measurements cannot be completely blinded to clinical findings, as he/she will perform the scans at certain time intervals, depending on the labour stage, that is established by the clinician’s cervical dilatation assessment. However, the sonographer will only record the images. The clinician will note the observations on a partogram-like sheet that is not available for the sonographer, who in turn, will perform the measurements offline, after birth.

Labour characteristics will be recorded by the clinician on the study datasheet: mode and time of delivery, neonatal Apgar score and birth weight, whether labour was spontaneous or induced, use of oxytocin or epidural

Figure 1 Implementation of the SonoLabor Study. DA, direction angle; FHPo, fetal head position; FHPr, fetal head progression; FHRo, fetal head rotation; HPD, head to perineum distance; MLA, midline angle; PA, progression angle; PD, progression distance; US, ultrasound.

(PD), head direction angle (DA) and head to perineum distance (HPD).

Caput measurement, if present.

Moulding notation, if present.
anaesthesia, occipital position at delivery. Maternal characteristics will be retrieved from the hospital records (patient files) by the personnel involved in data centralisation: age, height, weight, ethnicity, parity, gestational age.

The US labour assessments should not be biased by confounders that influence the quality of the clinical evaluations in labour (obesity, anterior placenta, caput, moulding), because the visualisation of the fetal skull and pubic symphysis is easily achievable even in such conditions. To ensure protocol fidelity, all sonographers will have completed a 1-day workshop and will participate in group supervision sessions programmed weekly in the first month of the study. This approach proved to be successful during the previous pilot study conducted in our centre.

The information provided to the clinician regarding the US determination of the FHPo in case of clinical–sonographic discordance before instrumental or operative delivery could represent a theoretical bias of the study. However, many tertiary centres already use the US determination of the FHPo in such situations, and this aspect does not interfere with the objectives of our study.

### Outcome measures

#### Primary outcome

The primary objective of this study is the elaboration of nomograms for the longitudinal US assessment of labour in unselected nulliparous and multiparous women with fetal cephalic presentation. The nomograms represent the evolution of the progression markers (PA, DA, PD, HPD) in relation to time and cervical dilatation.
Secondary outcomes

- To compare the US pattern of labour evolution in nulliparous and multiparous women.
- To study the influence of occiput position, BMI and parturient age on the mechanism of delivery evaluated by US.
- To compare the labour clinical trend from our study data with the Friedman studies\textsuperscript{35,36} and other recent research on the partogram.\textsuperscript{39}
- To correlate the US findings with classic clinical estimations:
  - Correlation of the FHPo determined by US with FHPo clinically estimated (by digital vaginal evaluation (VE)).
  - Correlations between the US FHPr parameters and between US and clinical (head station) FHPr parameters.
  - The concordance between the fetal head station evaluations derived from US measurements and clinical digital estimations.
- To investigate the correlations between the data of the participating centres.
- To analyse the evolution of the sonographic measurements in spontaneous vaginal delivery versus obstructed labour cases, in nulliparae versus multiparae, and in occiput anterior deliveries versus those with persistent occiput posterior.
- To evaluate the capability of the US technique to predict the labour outcome (vaginal or caesarean birth) in both nulliparous and multiparous women.

Data collection and management, and quality control

To ensure protocol fidelity, the sonographers will have completed a 1-day workshop and will participate in group supervision sessions, programmed weekly in the first month of the study. This approach proved to be successful during the previous pilot study conducted in our centre. The following procedures are to be followed:

- At the initial workshop, the standardised procedures regarding data collection, encoding of the clinical and US data and electronic storage will be established.
- The principal investigators will organise training sessions to provide instructions on the protocol and study procedures for the sonographers at the beginning of the study.
- Monthly meetings will take place between study site personnel to discuss issues related to the conduct of the study and supplementary convocations will be announced whenever necessary.
The principal investigators in the three centres will be available for consultation by telephone at request.

Interim analyses monthly—the data manager will evaluate the data with the statistics personnel and will conduct a quality review of the database. The results of the interim analyses will be discussed between the principal investigators, who decide whether to continue, stop or modify the trial.

All the collected data will be anonymised. The data will be collected by the research team, processed and stored in the www.zenodo.org research depostitory.

**Statistical methods**

**Sample size estimation**

Although several studies have investigated the clinical course of labour, until present we do not have data regarding the nomograms for US evolution of labour. The number of patients enrolled in the clinical partogram studies varies widely. However, the Friedman’s study that still serves as the basis of how most physicians define normal labour enrolled 500 nulliparous women at term.37 38 On the other hand, we have a recent large, but retrospective study that analysed the clinical labour records of more than 62 000 women from 19 hospitals across the USA and concluded that these criteria created 50 years ago may no longer be applicable to contemporary obstetric populations and for current obstetric management.39

Regarding imaging studies, the prospective research on the trend of the labour progress using intrapartum transperineal US gathered less than 100 cases each.34 36 44

The primary outcome of our study will be centile charts for each US progression marker in relation to time.

An important challenge of our study is to achieve a sufficient number of OP cases in both nulliparous and...
Multiparous women. According to the Central Limit Theorem and the Large Enough Sample Condition, a sample size of at least 30 items is sufficient for describing a ‘normal’ behaviour of the sample, even if it is not governed by the Gaussian distribution. By looking at the \( t \) table, we can see that when using around 30 df, the value of \( t \) becomes approximately equal to the value of the \( z \) statistics.\(^{45}\) Taking into account previously published studies, the overall rate of occiput posterior deliveries in nulliparous is around 7.2\%, whereas for the multiparous deliveries is around 4\%. Only in 65\% of these cases the outcome is vaginal birth. This implies that the corresponding sample size is 642 nulliparous women, and 1154 multiparous women who give their consent to participate in the study. Using this sample size, we achieve a suitable statistical power two-type of null hypothesis with default statistical power goals \( p \geq 0.95 \) and type I error \( \alpha = 0.05 \) level of significance.

In our pilot study, a cervical dilatation of more than 4 cm was noted in 16.34\% of the nulliparous women and 37.5\% of the multiparous women who were admitted to the hospital with labour criteria. In such cases, data from the beginning of labour will not be available for calculation. In order to achieve the sample mentioned above with patients registered from the beginning of labour, we adjusted the study size to include 767 nulliparous women and 1846 multiparous women.

Statistical analysis
The statistical analyses will be performed by IBM SPSS Statistics for Windows, V.22.0. (IBM Corp).

Descriptive statistics will be produced for all study variables (mother’s age, height, weight, parity, gestational age, mode and time of delivery, whether labour was spontaneous or induced, use of oxytocin or epidural anaesthesia, occiput position, PA, PD, HDA, HPD). Continuous
variables will be presented as the mean and SD or median, if appropriate. Categorical data will be presented as frequency and percentage.

Data will be first tested for normality and equal variance. Clinical obtained data from our study will be compared with similar data from other partograms. The results between groups (maternal, labour and neonatal characteristics of women assessed by classic clinical partograms or a more recent partogram and our sonopartogram) will be compared using \( \chi^2 \) test or Fisher’s exact test (for categorical variables), and Student’s t-test or Mann-Whitney test where applicable (for continuous variables) with a statistical significance level set at \( p<0.05 \).

We will analyse the agreement between sonopartogram and clinical partogram in estimating FHPo and fetal head station. For the FHPo, we will assess the level of agreement between US and digital VE using Cohen’s kappa statistics. Correlation coefficient (Pearson’s correlation or the Spearman’s rank correlation or Kendall’s rank correlation if appropriate) and linear regression will be employed for analysing the strength of association between the fetal head station estimated by digital VE and the US parameters (HPD, PA, PD and HDA).

Pearson’s correlation and regressions will be used for the evaluation of correlation between US parameters (PA, PD, HDA and HPD) and between US and time to delivery and digital VE (head station) for various clinical situations (nulliparous and multiparous, fetuses with OA and those with persistent OP position).

Reference ranges (90% range between 5th and 95th centiles) and the 95% CI will be constructed for each US parameter, and evolution in time will be displayed in graphic form separately for nulliparous and for multiparous. Predictive ability of each US parameter for vaginal delivery will be assessed by calculating sensitivity, specificity, positive predictive value, negative predictive value and likelihood ratio and by plotting receiver operating characteristic (ROC) curve.

In order to identify factors that predict vaginal birth, for each subgroup population (nulliparous and multiparous), all analyses will use appropriate (that is, logistic or linear) regression models, with results presented as point estimates (ORs or difference in means), 95% CIs and \( p \) values. Further secondary analyses will involve planned subgroup analyses and will use multivariable regression models. In all models, predictors (like maternal age, gestational age, clinically assessed cervical dilatation, maternal BMI) will be selected for inclusion in regression. We plan to include in our model covariates such as HPD, PA, PD, HDA and OP position. Based on the probabilities predicted by the logistic models, ROC curves will be constructed and we will calculate and report the area under the curve, sensitivity and specificity rates with 95% CI in predicting vaginal mode of delivery.

The time from the US examination at the beginning of active phase of labour to vaginal delivery will be evaluated with Kaplan-Meier and Cox regression analysis. Data for women with caesarean section will be censored. In the Cox regression analyses, FHPo and fetal head station parameters will be tested as possible predictive factors. In additional analyses, we will adjust for maternal age, BMI, gestational age and parity as possible confounders.

**Reporting of adverse events**

Prenatal ultrasonography appears to be a safe investigation method, as until today there has been no study reported suggesting otherwise (statement approved by the International Society of Ultrasound in Obstetrics and Gynecology Board in September 2011 and by the World Federation of Ultrasound in Medicine and Biology Council in August 2011). US is routinely used in everyday clinical practice for assessment of neonates,
including cranial and cerebral examination. However, US involves energy exposure and that requires further investigation.

Regarding the perception of labouring women about US, there have been no reports in the literature of US causing discomfort.

All adverse events reported spontaneously by patients or observed by the obstetricians will be recorded. When an adverse event occurs, the treating physician will take all necessary and appropriate measures to ensure the safety of the patient.

**Ethical considerations and dissemination**

**Ethics approval and consent to participate**

Ethics approval of the study protocol was obtained from the ethics committees of the universities in the three centres. The trial is approved by the University of Medicine and Pharmacy of Craiova Committee of Ethics and Academic and Scientific Deontology (No: 18/26.02.2016).

**Informed consent**

The US operator on duty will be responsible for explaining the procedure to the participants and for obtaining a written informed consent from all women accepting to take part in the study.

Regarding the unforeseen complications or health damage that may occur during or after labour, the management of labour and delivery is made exclusively based on the traditional clinical evaluation, by senior physicians. The US study protocol is only observational, without any obstruction for the clinical manoeuvres. The only potential sonographic intervention in the clinical assessment of labour is due to the ethical issues regarding the neonatal outcome when instrumental delivery is attempted. Thus, the attending obstetrician will be informed in case of clinical and US discordance.

On the other hand, it is made clear to all participants that US is considered safe in the third trimester and after birth, both for the mother and the baby.

**Compensation and insurance for harmed patients**

There will be no special financial compensation; however, any negligence on the part of the physician may be covered by the doctor’s liability insurance.

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**Contributors** DGI, PA and ST conceived and designed the study and will supervise the study implementation. DGI drafted the protocol of the study. LMD, GD, MLC, RD, CP, LZ, MN and DT revised and refined the study protocol for an optimal implementation and provided final approval of the version to be published. MLC, SB and RS provided methodological and statistical expertise and will conduct the statistical analysis. DGI and RD drafted a PhD salary grant proposal for a previous approved pilot study. GD, RD, CP, LZ, RN, DR and MF will be responsible for study management, staff training and supervision, and data centralisation. DGI, PA and DT are the directors of the sites and they will provide clinical expertise and on-site management of the study. All authors critically reviewed and approved the final version of the manuscript.

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