

Contribution of ultrasonography to the prediction of the induction-delivery interval: The ECOLDIA prospective multicenter cohort study

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Title

Contribution of ultrasonography to the prediction of the induction-delivery interval: The ECOLDIA prospective multicenter cohort study.

Authors

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1 Highlights:

Previous studies reported contradictory results that might be explained
 by heterogeneous populations

We sought to evaluate a large population of women homogeneous for
 cervical status using a robust methodology. Thus, bishop score was an
 original inclusion criterion when selecting a population to study the
 prediction of induction-delivery interval.

- It is the largest cohort of women to evaluate the induction-delivery
 interval.
- Moreover, women undergoing induction of labor with a harmonized
 single-agent protocol in order to reduce heterogeneity of study
 population.
- Transvaginal ultrasound cervical length can significantly predict
 induction-delivery interval with parity.
- 15

16 **ABSTRACT**

Introduction: To evaluate the ability of preinduction ultrasonographic cervical
length to predict the interval between induction and delivery in women at term
with a Bishop score of 4 to 6 at induction.

Study design: This multicenter prospective observational cohort recruited 334 women from April 2010 to March 2014. Inclusion criteria were women with singleton pregnancies at a gestational age ≥37 weeks, with no previous caesarean, a medical indication for induction of labor, and a Bishop score of 4, 5, or 6. All women underwent cervical assessment by both transvaginal ultrasound and digital examination (Bishop score). The induction protocol was
standardized. The primary outcome measure was the induction-delivery interval.
Hazard ratios (HR) and their 95% confidence intervals (95% CI) were used to
assess potential predictors.

Results: Mean gestational age at induction was 40.1 weeks, 60.8% of the women were nulliparous, and the cesarean rate was 13.4%. The mean induction-delivery interval was 20.8 h (\pm 10.6). Delivery occurred within 24 h for 56.9% (n=190) of the women. An ultrasonographic cervical length measurement less than 25 mm (HR=1.50, 95% CI 1.18–1.91, P<0.01) and parity (HR=1.41, 95% CI 1.21–1.65, P<0.01) appeared to predict induction-delivery interval. The cervical length cutoff to reduce the induction-delivery interval was 25 mm.

36 Conclusion: A cervical length cutoff of 25 mm was associated with shorter 37 induction-delivery interval in women at term with a Bishop score of 4 to 6.

38

39 **Abbreviations**

40 TVUS: transvaginal ultrasonography

41

42 **INTRODUCTION**

Induction of labor occurs in nearly 22% of pregnancies (1). An important challenge in induction of labor is predicting which patients will have vaginal deliveries and the time interval from induction to delivery. We believe that information about delay is an important clinical feature that can affect women's information and satisfaction when inducing labor. For this reason, we were focused on delay rather than outcome of the delivery. The Bishop score is 49 currently the standard method for evaluating local cervical ripening before 50 induction (2). A Bishop score of six or more is considered favorable for induction 51 of labor, so that oxytocin can be recommended to start induction (3). An 52 intermediate Bishop score is interpreted to mean that the cervix is unfavorable, in which case it is recommended that women undergo cervical ripening before 53 54 oxytocin administration. Their management thus relies on digital cervical 55 examination, known to have a subjective measurement with high inter- and 56 intra-observer variability and a poor predictive value for delivery outcome (4). It 57 may thus be of limited value, especially in women with intermediate Bishop 58 scores (4 to 6). Accordingly, other types of preinduction cervical evaluations 59 have been suggested, such as ultrasound assessment, because they might be reproducible and more objective as well as more acceptable to women (5). In 60 61 women with a low Bishop score, it could also be a useful tool for predicting time 62 to delivery.

Previous studies comparing the Bishop score with transvaginal ultrasonography 63 64 (TVUS) of the cervix to predict time to delivery or delivery outcome have reported contradictory results (6-17). Their differences might be explained by 65 small samples (between 43 to 266 women included) and heterogeneous 66 67 populations that further differ for their Bishop scores at inclusion, gestational age at induction, and main outcome. Moreover, induction methods were not 68 69 standardized within studies and differed between them. Similarly. 70 ultrasonographic measurements were not homogeneous and various 71 parameters were evaluated: cervical length, width, dilatation, posterior cervical 72 angle, and lower segment thickness. We sought to evaluate a large population

of women homogeneous for cervical status and undergoing induction of laborwith a harmonized single-agent protocol.

Accordingly, the aim of this study was to evaluate if preinduction ultrasonographic cervical measurements is associated with the inductiondelivery interval in women at term.

78

79 MATERIALS AND METHODS

80 The prospective multicenter cohort study ECOLDIA (Echographie du COL dans 81 l'évaluation du Délai Induction Accouchement) took place from April 2010 82 through March 2014, in three tertiary hospital centers in France. Screening took 83 place among all women for who induction of delivery was indicated whatever 84 the indication. The inclusion criteria required that women be pregnant with a 85 singleton live fetus in cephalic presentation at 37 to 42 weeks of gestation, and 86 have a medical indication for labor induction, a Bishop score of 4 to 6 during the hour before induction, and no contraindication for dinoprostone. The Bishop 87 88 score was assessed by digital examination by a midwife and was calculated according to the position, consistency, shortening and dilation of the maternal 89 90 cervix and the station of the fetal presenting part (2). The exclusion criteria 91 were: a previous cesarean delivery, any indication for an elective cesarean, 92 cervical cerclage for this pregnancy, congenital uterine malformation, a history 93 of uterine surgery, fetal abnormality, or an age younger than 18 years.

The studied factor was the ultrasonographic cervical length assessed one hour before induction. TVUS was performed in all cases by an obstetrician blinded to the results of the clinical examination. All participants were blinded to the

97 cervical length. A junior or senior ultrasound certified obstetrician used a 5-9 98 MHz transvaginal probe and a Voluson E8 (GE Healthcare, Milwaukee, WI, 99 USA) for the TVUS. A standardized procedure was used for cervical 100 measurement: the woman had to have an empty bladder and was placed in a 101 supine position with legs abducted. The operator placed the probe in the 102 anterior vaginal fornix to obtain a sagittal view of the cervix, avoiding undue 103 pressure on it to avoid false elongation of the images. He or she then identified 104 the internal and external os, using the endocervical mucosa to define the level 105 of the internal os, and checked for and noted any funneling at the internal os. 106 The cervical canal was magnified to obtain at least 75% of the image. Three 107 measurements of the distance between the internal and external os were taken 108 over a period of about 3 minutes, and the shortest measurement of the cervical 109 length was recorded. Width and dilatation of internal os were also evaluated.

The following data were also collected: age, body mass index, parity, term of pregnancy, weight gain, indication for induction, analgesia during labor, oxytocin, mode of delivery and, if a cesarean was performed, its indication, interval from the start of induction to delivery, Apgar score, arterial pH, and neonatal hospitalization in the intensive care unit.

115 Cervical ripening for all women was performed with the vaginal prostaglandin 116 E2 slow-release system (Propess® 10 mg, Ferring, Gentilly, France), which 117 releases the medication at 0.3 mg/h, for 24 hours. If labor had not started 24 118 hours after cervical ripening began, it was induced by artificial rupture of 119 membranes and oxytocin administration, in accordance with French guidelines 120 (18). 121 The primary outcome was the induction-delivery interval, defined as the period122 from the start of cervical ripening to delivery in hours.

123 Statistical analyses

124 It was difficult to formulate a quantitative hypothesis regarding the association 125 between ultrasound variables and induction-delivery time because there were 126 no data available in the literature. Based upon the rates of induction in the 127 participating units, our aim was to include 400 women to be able to conduct the 128 study with sufficient power and reasonable precision.

129 Continuous data are presented as medians with their interguartile ranges (1st 130 quartile-3rd quartile) or means and their standard deviations, and categorical 131 data as counts and percentages (normal distribution was verified by histograms). 132 Receiver operating characteristic (ROC) curves were generated to identify the 133 best cutoff values with maximum efficiency for TVUS cervical length as 134 categorical predictors of the induction-delivery interval. To construct the ROC 135 curve, the binary outcome was the delivery within 24 hours. We conducted 136 survival analysis of the data with the Kaplan–Meier method and the log rank 137 test.

Then, to take into account the competing risks between vaginal delivery and cesarean delivery occurs during labor, we used a Fine and Gray regression model. Indeed, a woman who had a C-section because she fails to progress at 6cm for example will delivery faster than a woman who starts labor quickly and had a vaginal delivery. The associations between the ultrasound measurement (functional cervical length and funnel width), parity, maternal age, gestational age at induction, BMI, indication for induction (PPROM versus other indications) and induction-delivery interval are presented with hazard ratios (HR) and 95%
confidence intervals (95% CI).

Analyses were conducted with R version 3.1.3. Differences were defined assignificant when P<0.05.

149

150 Ethical approval

151 Written informed consent was obtained from all participants. The ethics 152 committee approved this study. This clinical trial was registered as 153 NCT02570620. The study follows the STROBE statement guidelines for 154 reporting observational studies (19).

155

156 **RESULTS**

This study included 342 women and analyzed 334 (Figure 1); their characteristics and outcomes are shown in Table 1. Regarding neonatal outcomes, the median Apgar score was 9.7 (\pm 0.7), the arterial pH was 7.2 (\pm 0.08), and the venous pH was 7.3 (\pm 0.07). There was no difference in perinatal outcomes between women who delivered before or after 24 hours (p=0.846).

Among primiparous women, 43.8% (n=89) gave birth within 24 hours. The average functional cervical length before induction was 18 mm (\pm 11 mm) for women who delivered before 24 hours and 24 mm (\pm 12mm) for those who delivered with an interval of more than 24 hours. The results of ultrasound cervical length measurement compared with clinical assessment are shown in Table 2. There was a good correlation between the two measurements. Among women who gave birth within >24 hours, 96.5% received oxytocine after 169 dinoprostone. The mean induction-delivery interval for the entire population was 170 20.8 hours (\pm 10.6). Delivery occurred within 24 hours for 56.9% of the women 171 (n=190) with a mean induction-delivery interval of 13.3 hours. For the remaining 172 43.1% of women (n=144) who gave birth more than 24 hours after induction 173 began, the mean induction-delivery interval was 31.2 (±7.6) hours. This mean interval was 20.5 hours (± 10.6) for women who had a vaginal delivery, 25.4 174 175 hours (± 10.2) for those with a cesarean delivery (P=0.004; reference group 176 "vaginal delivery"), and 31.2 hours (±8.9) for the cesareans indicated for "failure 177 to progress" or "failed induction".

178 First, we evaluated the parity and TVUS cervical length to predict the induction-179 delivery interval. Cutoff values for cervical length were obtained from the ROC curve (Figure 2). The AUC of the ROC curve was 0.65 for predicting induction-180 181 delivery interval based on functional cervical length. We found no difference between a Bishop score of 4, 5, or 6 and the induction-delivery interval 182 183 (P=0.119). This result confirms the homogeneity of the study population. A 184 cervical length less than 25 mm at induction was associated with a shorter 185 induction-delivery interval (Log rank test, P<0.001) (Figure 3). We evaluated 186 some others ultrasonographic measurements such as width and dilatation of 187 internal os. Multiparity was also associated with a shorter induction-delivery 188 interval (Figure 4).

189 Second, we evaluated the induction-delivery interval in a multivariable 190 regression model, taking cesarean delivery into account as a competing risk, 191 adjusted for TVUS cervical length, funnel width, parity, maternal age, 192 gestational age at induction, BMI and indication for induction (PPROM versus other indications). The cervical length and funnel width as measured by
ultrasonographically, parity and induction for PPROM were the factors that were
significant predictors of the induction-delivery interval (respectively, HR=1.32,
95% CI 1.03–1.69, P=0.03; 1.02 [1.01;1.04] p=0.01; HR=1.50, 95% CI 1.27–
1.77, P<0.01; 2.02 [1.43;2.86] p<0.01) (Table 3).

198

199 COMMENT

Our results showed that measuring cervical length by ultrasound before induction improved the prediction of the induction-delivery interval in a large prospective cohort of women at term with a Bishop score of 4–6. The cervical length cutoff associated with a shorter induction-delivery interval was 25 mm. Funnel length was also associated with a reduced delay. Multiparity and PPROM were also predictive of a shorter induction-delivery interval.

These results are consistent with data from the literature, which find a significant association between the ultrasonographic measurement of the cervix and the induction-delivery interval, or the success of labor induction (6–12). Other studies, however, have not found that cervical ultrasound is more useful than the Bishop score for predicting time to delivery (13–17). Conflicting data from the literature can be explained by numerous biases in the studies.

First, the numbers included are generally small. Among the seven studies that calculated the required sample size and included more than 100 women, three showed that the Bishop score was more useful than TVUS. The main outcome criteria differed and were difficult to compare. Few studies considered the Bishop score (which was not initially created to predict issues of induction) as an inclusion criterion, as we did. In addition, the populations are heterogeneous
for gestational age, parity, and method of induction. All of this could lead to an
underestimation of the value of TVUS in decisions about labor induction.

220 A meta-analysis including two small randomized controlled trials of 234 women 221 that compared the Bishop score and TVUS for assessing preinduction cervical 222 ripening (20) did not demonstrate that either method was superior to the other 223 for determining the induction-delivery interval. In the first trial, this interval was 224 11.2 hours (IQR 7.8 to 15.9) for the Bishop score arm versus 9.5 hours (IQR 5.6 225 to 14.7) in the TVUS arm (21). In the second randomized controlled trial, the 226 median induction-delivery interval reported was 10.3 hours (95% CI 7.0-13.5) in 227 the Bishop score group, and 10.9 hours (95% CI 9.4-12.3) in the TVUS group 228 (22). These differences were not significant. The median interval in the two 229 randomized controlled trials was half of our interval (i.e., a mean induction-230 delivery interval of 20.5 hours (SD 10.6)), likely explained by the heterogeneous 231 populations in those trials; 20% of the women had a Bishop score greater than 232 6. Our study included only women with a Bishop score of 4, 5, or 6; they were 233 both nulliparous and parous, and received prostaglandins for no more than 24 234 hours before oxytocin infusion.

On the other hand, these two trials reported that cervical length cutoff values of 30 mm (21) and 28 mm by TVUS could reduce the need for intracervical prostaglandin treatment by 35% and 50% respectively, without affecting the success rate for induction. Thus, TVUS might change practices by identifying women who despite an intermediate cervix can undergo oxytocin administration to avoid unnecessary exposure to cervical ripening by prostaglandins. That is 241 the reason why the trial was designed to evaluate the delay and none the 242 success of the delivery as a primary outcome. In our population, the cutoff to 243 predict a shorter induction-delivery interval was 25 mm. Our results were 244 consistent those of with other studies: Gabriel et al. found that women with an unfavorable Bishop score had a shorter duration of labor when their cervical 245 246 length was less than 26 mm (8). A cutoff value at 28 or 30 mm may more precisely define a Bishop score less than 4 (21,22). The transvaginal cervical 247 248 length measurement may be superior to the Bishop score in evaluating the 249 ripened cervix. Several parameters TVUS could thus be useful supplementary 250 information for obstetricians to evaluate the benefit of cervical ripening.

251 One strength of this study is its homogeneous population, defined by an 252 inclusion criterion based on the Bishop score, intended to limit selection bias. 253 Another is the standardization of the induction protocol for all women, and the 254 large size of the cohort. The study's principal limitation is its observational 255 nature. The cutoff value of sonographically measured cervical length to 256 determine the value of cervical ripening must be validated.

257

258 CONCLUSION

A cervical length cutoff of 25 mm may help clinicians to predict the inductiondelivery interval in women with an unripe cervix. We postulate that knowledge of this interval is a useful tool for clinicians to allow them to adapt management to the degree of urgency in obstetric care and to economic considerations. Further studies are needed to assess whether cervical ripening is necessary in this population.

- 265 Conflicts of interest
- 266 None

267

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271

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- 275

276 **References**

- Blanc-Petitjean P, Salomé M, Dupont C, Crenn-Hebert C, Gaudineau A, Perrotte F, et al. État des lieux des pratiques de déclenchement en France.
 Gynécologie Obstétrique Fertil Sénologie. 2019 Jul;47(7–8):555–61.
- Bishop EH. PELVIC SCORING FOR ELECTIVE INDUCTION. Obstet
 Gynecol. 1964 Aug;24:266–8.
- Eggebø TM, Økland I, Heien C, Gjessing LK, Romundstad P, Salvesen
 KÅ. Can ultrasound measurements replace digitally assessed elements of
 the Bishop score? Acta Obstet Gynecol Scand. 2009 Jan;88(3):325–31.
- 4. Hendrix NW, Chauhan SP, Morrison JC, Magann EF, Martin JN, Devoe
 LD. Bishop Score: A Poor Diagnostic Test To Predict Failed Induction
 Versus Vaginal Delivery: South Med J. 1998 Mar;91(3):248–52.
- 5. Tan PC, Vallikkannu N, Suguna S, Quek KF, Hassan J. Transvaginal
 sonographic measurement of cervical length vs. Bishop score in labor
 induction at term: tolerability and prediction of Cesarean delivery.
 Ultrasound Obstet Gynecol. 2007 May;29(5):568–73.
- Boozarjomehri F, Timor-Tritsch I, Chao CR, Fox HE. Transvaginal ultrasonographic evaluation of the cervix before labor: Presence of cervical wedging is associated with shorter duration of induced labor. Am J Obstet Gynecol. 1994 Oct;171(4):1081–7.

- Daskalakis G, Thomakos N, Hatziioannou L, Mesogitis S, Papantoniou N,
 Antsaklis A. Sonographic Cervical Length Measurement before Labor
 Induction in Term Nulliparous Women. Fetal Diagn Ther. 2006;21(1):34–8.
- 8. Gabriel R, Darnaud T, Chalot F, Gonzalez N, Leymarie F, Quereux C.
 Transvaginal sonography of the uterine cervix prior to labor induction:
 Uterine cervix prior to labor induction. Ultrasound Obstet Gynecol. 2002
 Mar;19(3):254–7.
- Pandis GK, Papageorghiou AT, Ramanathan VG, Thompson MO,
 Nicolaides KH. Preinduction sonographic measurement of cervical length
 in the prediction of successful induction of labor: Cervical length and
 successful induction. Ultrasound Obstet Gynecol. 2001 Dec;18(6):623–8.
- Rane SM, Guirgis RR, Higgins B, Nicolaides KH. The value of ultrasound
 in the prediction of successful induction of labor: Ultrasound and induction
 of labor. Ultrasound Obstet Gynecol. 2004 Oct;24(5):538–49.
- Verhoeven CJM, Opmeer BC, Oei SG, Latour V, van der Post JAM, Mol
 BWJ. Transvaginal sonographic assessment of cervical length and
 wedging for predicting outcome of labor induction at term: a systematic
 review and meta-analysis: Cervical length in induction of labor. Ultrasound
 Obstet Gynecol. 2013 Nov;42(5):500–8.
- Ware V, Raynor BD. Transvaginal ultrasonographic cervical measurement
 as a predictor of successful labor induction. Am J Obstet Gynecol. 2000
 May;182(5):1030–2.
- 318 13. Jwatson W, Stevens D, Welter S, Day D. Factors predicting successful
 319 labor induction. Obstet Gynecol. 1996 Dec;88(6):990–2.
- 14. Chandra S. Transvaginal ultrasound and digital examination in predicting
 successful labor induction. Obstet Gynecol. 2001 Jul;98(1):2–6.
- 322 15. Gonen R, Degani S, Ron A. Prediction of successful induction of labor:
 323 comparison of transvaginal ultrasonography and the Bishop score. Eur J
 324 Ultrasound. 1998 Aug;7(3):183–7.
- Rozenberg P, Chevret S, Chastang C, Ville Y. Comparison of digital and
 ultrasonographic examination of the cervix in predicting time interval from
 induction to delivery in women with a low Bishop score. BJOG Int J Obstet
 Gynaecol. 2005 Feb;112(2):192–6.
- Rozenberg P, Goffinet F, Hessabi M. Comparison of the Bishop score,
 ultrasonographically measured cervical length, and fetal fibronectin assay
 in predicting time until delivery and type of delivery at term. Am J Obstet
 Gynecol. 2000 Jan;182(1):108–13.
- 18. HAs. Les recommandations pour la pratique clinique. Déclenchement
 artificiel du travail à partir de 37 semaines d'aménorrhée. Available:

- https://www.has-sante.fr/jcms/c_666473/fr/declenchement-artificiel-du travail-à-partir-de-37-semaines-d'aménorrhée.
- 19. Cuschieri S. The STROBE guidelines. Saudi J Anaesth. 2019;13(5):31.
- 20. Ezebialu IU, Eke AC, Eleje GU, Nwachukwu CE. Methods for assessing
 pre-induction cervical ripening. Cochrane Pregnancy and Childbirth Group,
 editor. Cochrane Database Syst Rev [Internet]. 2015 Jun 12 [cited 2020
 Feb 14]; Available from:
 http://doi.wiley.com/10.1002/14651858.CD010762.pub2
- Bartha JL, Romero-Carmona R, Martínez-del-Fresno P, Comino-Delgado
 R. Bishop score and transvaginal ultrasound for preinduction cervical
 assessment: a randomized clinical trial: Bishop score and TVS for
 preinduction cervical assessment. Ultrasound Obstet Gynecol. 2005
 Feb;25(2):155–9.
- Park KH, Kim SN, Lee SY, Jeong EH, Jung HJ, Oh KJ. Comparison
 between sonographic cervical length and Bishop score in preinduction
 cervical assessment: a randomized trial. Ultrasound Obstet Gynecol. 2011
 Aug;38(2):198–204.
- 352
- 353
- 354
- 355

356 Legends of figures and tables

- 357 Figure 1: Flow chart of the study
- 358 Figure 2: ROC curve of cervical length as a function of induction-delivery
- 359 interval (< or >24h).
- 360 Figure 3: Kaplan-Meier survival curves of induction-delivery interval according
- 361 to TVUS cervical length measurement before induction. The solid line
- 362 represents a cervical length < 25 mm. The dotted line represents a cervical
- length \geq 25 mm. Log rank test p<0.01.

Figure 4: Kaplan-Meier survival curves of induction-delivery interval according to TVUS cervical length measurement before induction. The solid line represents nulliparous women. The dotted line represents parous women. Log rank test p<0.01.

368

369

370 Table 1: Characteristics of 334 women analyzed.

371 Table 2: Clinical cervical length in comparison to the cervical length measured

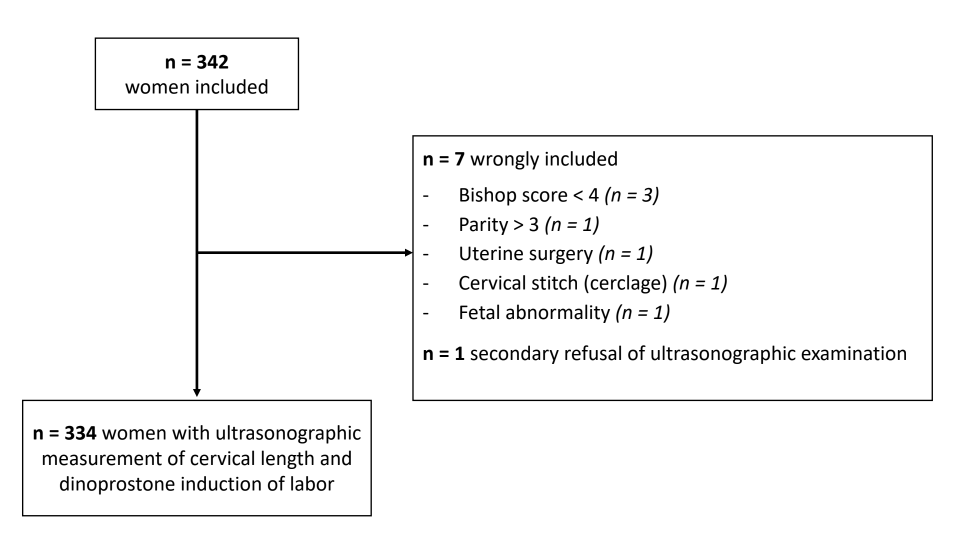
by transvaginal ultrasound without and with uterus pressure.

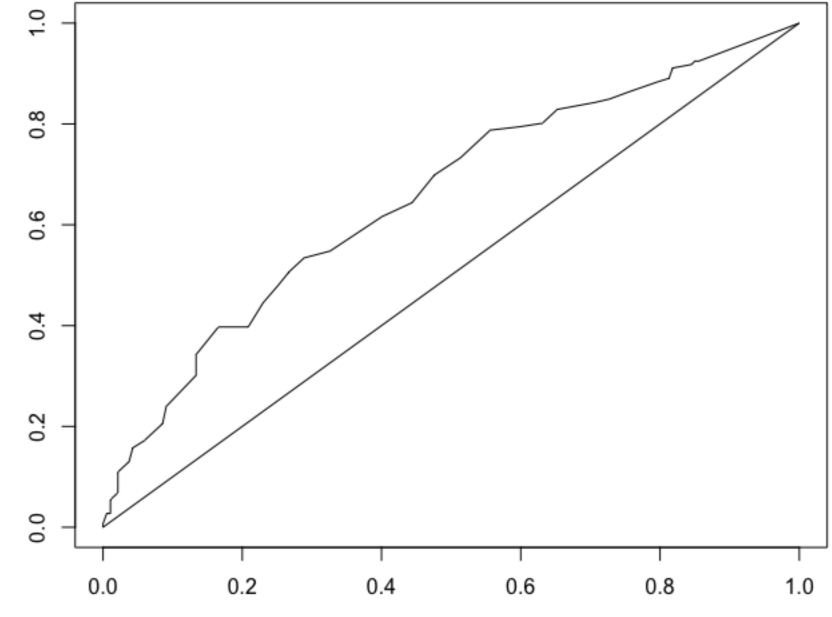
373 Table 3: Evaluation of induction-delivery interval in multivariable regression

374 model taking cesarean delivery into account as a competing risk, adjusted for

375 Bishop score, ultrasound cervical length, and parity

376

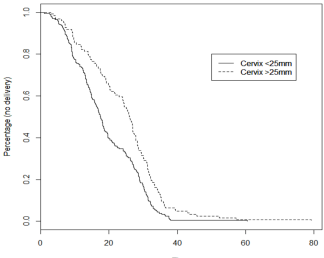




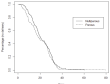
Cervical length as a function of induction-delivery interval (< or >24h)

False positive rate

True positive rate



Time



me

	Women (n=334)
Age (years)	29.0 (± 5.5)
Gestation at delivery (weeks' gestation)	40.1 (± 1.5)
BMI	25.5 (± 6.3)
BMI >30	37 (11.1)
BMI >35	29 (8.7)
Weight gain (kg)	13.7 (± 5.6)
Parity	
Nulliparous	203 (60.8)
Parous	131 (39.2)
Indication for induction of labor	
Prolonged pregnancy	87 (26)
Premature rupture of membranes	69 (20.6)
Gestational diabetes mellitus	44 (13.2)
Oligoamnios	37 (11)
Polyhydramnios	7 (2.1)
Decreased fetal movement activity	19 (5.6)
Maternal indication*	26 (7.8)
Fetal indication**	13 (3.9)
Hypertensive disease in	33 (9.9)
pregnancy, preeclampsia, or IUGR	
Bishop score	
4	133 (39.8)
5	146 (43.7)
6	55 (16.5)
Mode of delivery	
Spontaneous vaginal delivery	220 (65.9)
Instrumental delivery	59 (17.7)
Cesarean delivery	55 (16.5)
Ultrasonography	
Functional cervical length	20.4 (± 12.3)
Funnel length	4.7 (± 7.5)

Table 1: Means prenatal characteristics of 334 women analyzed.

Table 2: Clinical cervical length in comparison to the cervical length measured by

 transvaginal ultrasound without and with uterus pressure.

Cervical length measurement (in	Cervical length measurement (in	
mm), without uterus pressure	mm), with uterus pressure	
32 (± 8.5) [20-52]	30.9 (± 8.8) [21-53]	
29.3 (± 8.7) [9-54]	26.7 (± 8.5) [7-52]	
23.3 (± 8.1) [8-45]	21.1 (± 7.6) [7-41]	
19.5 (± 4.9) [14-25]	17.5 (± 3.1) [13-20]	
	mm), without uterus pressure 32 (± 8.5) [20-52] 29.3 (± 8.7) [9-54] 23.3 (± 8.1) [8-45]	

Mean ± satndard-deviation [*min-max*]

Table 3: Evaluation of induction-delivery interval in multivariable regression model taking cesarean delivery into account as a competing risk, adjusted for ultrasound measurement of cervical length and funnel width, and clinical characteristics that have been identified as potentially modifying the time to delivery.

Outcome	induction-delivery	P value
	interval HR (95% IC)	
Functional cervical length	1.32 [1.03;1.69]	0.03
Funnel length	1.02 [1.01;1.04]	0.01
Parity	1.50 [1.27;1.77]	<0.01
Maternal age	0.98 [0.96;1.01]	0.35
Gestational age at induction	1.05 [0.96;1.14]	0.25
BMI	0.98 [0.96;1]	0.14
Indication for induction PPROM or other	2.02 [1.43;2.86]	<0.01