ORIGINAL RESEARCH

Decision To Delivery Time and Its Predictors Among Mothers Who Underwent Emergency Cesarean Delivery At Selected Hospitals of Northwest Ethiopia, 2023: Prospective Cohort Study

Beker Ahmed Hussein¹, Beyene Sisay Damtew¹, Hinsermu Bayu Abdi¹, Temesgen Worku Gudayu²

¹Department of Midwifery, College of Health Sciences, Arsi University, Asella, Ethiopia; ²Department of Clinical Midwifery, College of Medicine and Health Sciences, University of Gondar, Gondar, Ethiopia

Correspondence: Beker Ahmed Hussein, Email bek50ahmed60@gmail.com

Background: The decision to delivery time is the interval between the decision and the childbirth by emergency caesarean delivery. The Royal College of Obstetricians and Gynecologists and the American College of Obstetricians and Gynecologists recommend that the decision to delivery time interval is less than 30 min. Additionally, the decision to delivery time varies across institutions and countries.

Objective: The aim of this study was to determine the decision to delivery time and its predictors among women who underwent emergency cesarean delivery at selected hospitals of Northwest Ethiopia, 2023.

Methods: An institutional-based prospective cohort study was conducted at selected hospitals of Northwest Ethiopia, among women who underwent emergency cesarean delivery from November 1 to January 30, 2023. A total of 285 participants were enrolled, and data collected using structured and pre-tested questionnaires. A systematic sampling technique was used. Data were entered into Epi-Data version 4.6 and then exported to STATA 15 for further analysis. The log rank test was utilized to compare group differences. The time is estimated by using the Kaplan–Meier curve and Cox proportional-hazard regression analysis was carried out to determine the predictors.

Results: From 285 participants, 56 (21.8%) women delivered within the recommended 30 min. The overall median survival time was undetermined and the restricted mean survival time was 48.9 min (95% CI: 47.4–50.5). The average decision to delivery time is affected among women who hesitate to accept consent (AHR: 0.17, 95% CI: 0.02–1.25), cord prolapses (AHR: 1.36, 95% CI: 0.46–3.94), rank of surgeon (AHR: 0.42. 95% CI: 0.42–1.08), no free operation room table (AHR: 0.27, 95% CI: 0.28–0.94), regional anesthesia (AHR: 0.56, 95% CI: 0.25–1.28), and use of a bladder flap (AHR: 0.33, 95% CI: 0.16–0.85).

Conclusion: Overall decision to delivery times among women who underwent emergency cesarean section at selected hospitals were longer than the recommended time.

Keywords: decision time, emergency cesarean delivery

Background

The period from the decision to perform a cesarean delivery until the baby was delivered is known as the decision to delivery time, and is measured in minutes.¹

The amount of time it takes to prepare the patient and the theater, provide anesthesia, and deliver the baby includes all of these factors. The decision to delivery time must be as quick as possible since an emergency caesarean section is done when the life of the mother or fetus is in urgent danger during delivery.² The suggested decision to delivery time is within 30 min, according to the American College of Obstetricians and Gynecologists (ACOG) and the Royal College of Obstetricians and Gynecologists (RCOG), as two such organisations.³ In contemporary obstetrics, normal caesarean

249

births are provided to women as an elective procedure for a several reasons, or they are carried out in an emergency due to fetal or maternal problems, or both.⁴ Caesarean sections can be categorized as emergency, urgent, planned, or elective depending on how serious the fetal or maternal condition is. The categorization states that emergency CS is carried out when the woman, the fetus, or both are in grave danger.⁵ The anesthesiologist and trained assistant, the obstetrician and assistant, a theater nurse, and neonatal support staff are all involved in the complicated and multidisciplinary cesarean section operation.⁶ It is necessary to gather this crew, as some of them must review the patient and also get her ready. Before surgery can be performed, certain patients may require additional care and stabilization to make sure their condition is at its best. Even while in the event of an emergency cesarean section the DDT must be regarded as having been completed as a goal, failure to stabilize the patient may increase the risk of maternal death and morbidity as a result of the surgery or anesthesia problem.⁷ A hospital that offers obstetric care should be able to react to an emergency within the required timeframe. The dangers associated with an emergency cesarean section, however, might be substantially higher in underdeveloped countries than in developed countries since it may take longer to start one there and resources may not be sufficient.⁸ Three delays prevent pregnant and laboring women from obtaining the professional care they require in a timely manner. These delays might be caused by a lack of supplies, inadequate knowledge or experience of the provider, or a lack of basic equipment.⁹

It has been noted that a variety of factors, such as team preparedness, team communication, team availability, severity of fetal-maternal problems, seniority of the surgeon, availability of obstetricians and anesthesia staff, time and day of operation, and kind of anesthesia, might impact the decision to delivery time (DDT).¹⁰ Facilities and service providers must be ready for quick emergency action and prepared in advance to solve the issue. To reduce morbidity and mortality associated with delayed decision to delivery time (DDT), further measures should be taken: a 24-hour cesarean surgery room that is housed within the labor and delivery ward, well-trained personnel who are available seven days a week, effective teamwork, and a delivery room with all the necessary equipment for emergency action.^{2,11} Compared to elective and vaginal births, the perinatal and maternal outcomes of emergency cesarean delivery are worse. Quality obstetric care services, such as reducing third delays and delays in getting appropriate treatment, such as shortening the time between decision to delivery interval is important for medico-legal problems as well as quality.^{12,13} Hence, this study aimed is to assess the decision to delivery time and its predictors among women who underwent emergency cesarean delivery.

Methodology

Study Area and Design

A prospective cohort study was conducted from November 1 to January 30, 2023 among women who underwent emergency cesarean delivery at selected hospitals of Northwest Ethiopia.

Source Population

All pregnant women who underwent emergency cesarean delivery at selected hospitals of Northwest Ethiopia were included.

Study Population

Pregnant women whose emergency cesarean delivery was done during the study period were included.

Eligibility Criteria

Inclusion Criteria

Pregnant women who underwent emergency cesarean delivery were included.

Exclusion Criteria

Women with multiple pregnancies were excluded.

Those stating total refusal of emergency cesarean delivery were excluded.

Sample Size Determination

A single population proportion formula was used to calculate the sample size by considering the following statistical assumptions.

P=the proportion of women who had a decision to delivery time interval below 30 min is 21.3%, from a study done in Ethiopia.¹⁴

 $Z\alpha/2$ =the corresponding Z score of the 95% CI.

d=margin of error (5%) and N=sample size.

 $N=(Z\alpha 2)^{2} \times p(1-p)/(d)^{2}$

 $N=(1.96)^2 \times 0.213 \times 0.787/(0.05)^2=259$, and then after adding 10% loss follow-up, the final sample size was 285.

Sampling Procedure

Two hospitals were selected by simple random sampling that were found in Northwest Ethiopia. Prior to data collection, the average number of women who underwent emergency cesarean delivery was estimated using reports from the previous three months. According to data from the past three months, each hospital had an average case flow of 562 for Debre Markos Comprehensive Specialized Hospital and 45 for Lumame Primary hospital. The sample size was allocated for two hospitals proportionally based on the estimated numbers of women who underwent emergency cesarean delivery in two hospitals over three months when the data collection was conducted. A kth interval $(562+45)/285\approx2$ of the total estimated numbers of women who underwent emergency cesarean delivery in the two hospitals over three months to the calculated sample size was approximated. The first participant (women) was recruited by a lottery method. The sample of women with emergency cesarean delivery was obtained from each hospital by using a systematic random sampling technique with a sampling interval of k=2 (Figure 1).

Study Variables

Dependent Variable

Occurrence of the decision to delivery time within 30 min.

Independent Variable

Sociodemographic:

age, educational status, marital status, place of residence, and attendant educational status.



Figure 1 Schematic presentation of the sampling procedure at selected hospitals of Northwest Ethiopia, 2023.

Obstetrics History

ANC follow-up, BPCR, presence of scar, and cervical dilation at decision.

Indication of Cesarean Section

Cord prolapses, NRFHBP, abruption, CPD, and uterine rupture.

Type of Anesthesia

General, spinal, and epidural.

Surgeon's Rank

Senior and resident.

Time and Day of the Week When the Surgery Will Be Performed

Weekday, weekend, daytime, and night-time.

Preparation for Emergency Cesarean Delivery

Resuscitative measures, obtaining consent, and unavailability of logistics.

Operational definitions

Event: if the delivery occurs in less or equal 30 min it was considered as an event, whereas above 30 min is considered as a censored observation.

Start time: decision time.

End time: delivery of the neonate by emergency cesarean section.

Emergency cesarean section: when the CS is done in labor or due to any complication that necessitates immediate cesarean delivery.

Decision time: when the surgeon who performed the procedure decided to do emergency cesarean delivery.

Data Processing and Analysis

Data were entered into Epi-Data version 4.2 after checking for completeness, and then the data were cleaned and transferred to STATA 15 statistical software for analysis. Before running the Cox proportional hazard regression model to simultaneously assess the association between multiple covariates and survival, multicollinearity was checked using the variance inflation factor (VIF) for continuous variables. All variables were found to be <10. The proportional hazard assumption (PHA) was checked to identify the combined effects of several covariates on the hazard ratio, using scaled Schoenfeld residual tests (PH test), and all variables fulfilled this assumption (P-value>0.05). The log rank test was utilized to compare group differences and the decision to delivery time was estimated using the Kaplan–Meier curve. To identify potential predictors of time to decision of delivery, a bivariable Cox proportional regression model was fitted for each explanatory variable and variables that would have a P-value less than 0.025 were included in the multiple Cox regression model. In the multiple Cox analysis, variables that have a P-value less than 0.05 were significant predictors of decision to delivery time. The adjusted hazard ratio was used to show the strength of association between each predictor variable and the outcome variable.

Data Collection Procedure

Data were collected using a standardized questionnaire. Six bachelor midwives were involved in the data collection. Once the study subject was recruited by systematic sampling techniques they were followed from the time the decision was made by the surgeon until the delivery of the neonate by emergency cesarean delivery. The decision to do an emergency cesarean delivery, the indication of the procedure, the time of the operation transfer, the duration of the anesthesia, the total time from the decision to the birth of the fetus, and the time the anesthesia team was informed were all obtained from the patient's charts and by direct observation. The moment the doctors made the decision to perform an emergency caesarean delivery and the time at which the operation was done were recorded. The length of time it took to

transfer to the operating room, choose anesthesia, administer it, and perform the procedure were all afterwards noted. afterward

Data Quality Control

Before data collection, a pretest on 5% of the sample was conducted. Both the supervisor and the data collectors received a two-day training. Daily completion and consistency checks were made on the completed questionnaire. Daily data cleaning was carried out, and the data collectors received quick feedback.

Results

Sociodemographic Characteristics of Study Participants

A total of 285 women who underwent category-1 cesarean section with a response rate of 100% were studied. The majority of women (120, 42.1%) in this study were aged between 25-29 years with a mean of 28 ± 5.006 . The majority of the respondents are married (261, 91.6%). Around 66 (23.2%) had a level of education of high school, diploma, and above. Around 135 (47.4%) are housewives. Regarding participants' residence, 194 (68.1%) were urban (Table 1).

Obstetric Characteristics of Study Participants

The findings showed that 37.9% of the respondents were primigravid. The majority of the respondents had ANC visits. Around 63.2% of women had been counseled on BPCR plan during ANC follow-up. Sixty-five (22.8%) participants had a previous history of cesarean section. The majority of the participants (258, 90.5%) accepted informed consent

Variables	Frequency (%)	Variables	Frequency (%)
Age in years		Educational status	
15–20	6 (2.1%)	Cannot read and write	51 (17.9%)
20–24 65 (22.8%)		Read and write	24 (8.4%)
25–29	120 (42.1%)	Elementary	55 (19.3%)
30–35	70 (24.6%)	High school	66 (23.2)
>35	24 (8.4%)	Preparatory	23 (8.1%)
Marital status		Diploma and above	66 (23.2%)
Single	7 (2.5%)	Residence	
Married	261 (91.6%)	Urban	194 (68.1%)
Divorced	12 (4.2%)	Rural	91 (31.9%)
Widowed	5 (1.8%)		
Occupational status			
Housewife	135 (47.4%)		
Government employee	48 (16.8%)		
Daily labor	13 (4.6%)		
Farmer	59 (20.7%)		
Merchant	29 (10.2%)		
Others	I (0.4%)		

Table ISocio-Demographic Characteristics of Women Undergoing Emergency CesareanDelivery at Selected Hospitals of Northwest Ethiopia, 2023 (n=285)

immediately after she was counseled for CS. In this study the main indication for emergency cesarean delivery is non-reassuring fetal heart rate (NRFHR) (Table 2).

Regarding the indication of emergency cesarean delivery, around 36.10% were women with NRFHR, 13% had grade three meconium, 10.9% with CPD, 10.20% are with two c/s scars, 4.60% are with cord prolapse, and 3.90% are with other indications (Figure 2).

Regarding the rank of the surgeon, from 285 study participants around 30.53% of emergency cesarean deliveries are performed by IESO, followed by MSc clinical Midwifery students who account for around 28.42%, and the remaining 25.26% and 15.79% are performed by senior and MSc clinical Midwife respectively (Figure 3).

The Incidence Rate and Median Decision to Delivery Time

The study participant was followed for a minimum of 18 min and a maximum of 56 min with a median follow-up time of 35 ± 8.14 min. Two hundred eighty five participants were followed for different periods of maximum of eighten minute and gives 10,060 persons minutes of obsrvation. Around 62 (21.8%) developed the event (delivered within 30 min) (95% CI: 17.2–27.3) while 223 (78.2%) women were censored (95% CI: 73–83.2). The overall decision to delivery time rate was 0.6 per 100-person minute (95% CI: 30–45.) The median decision to delivery time overall could not be calculated because the largest observed analysis time was censored.

Instead of computing the total median survival time, the mean survival time confined to the longest follow-up period was calculated. The overall restricted mean survival time was 48.9 min (95% CI: 47.4–50.5) (Figure 4).

Comparison of Survival Status for Different Factors with the Log Rank Test

The log rank test was performed to test the equality of survival curves for the presence of any notable variations in survival rates across the various levels of the categorical factors taken into account in the research. The test results in this study demonstrated a substantial difference in the survival function for several categorical variables. Accordingly, the Kaplan–Meier analysis revealed strong evidence of variations in decision to delivery timeframes. It was found that the mean decision to delivery time for women who accepted informed consent was 48.4 min (95% CI: 46.7–50.1), which

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Variables	Frequency (%)	Variables	Frequency (%)
Number of deliveries Primigravid Multigravida	108 (37.9%) 177 (62.1%)	Referral status Yes No	190 (66.7%) 95 (33.3%)
ANC follow-up Yes No	272 (95.4%) 13 (4.6%)	Previous history EC/D Yes No	65 (22.8%) 220 (77.2%)
BPCR Yes No	180 (63.2%) 105 (36.8%)	OR material prepared Yes No	247 (86.7%) 38 (13.3%)
Informed consent Agreed Hesitate	258 (90.5%) 27 (9.5%)	Free functional OR table Yes No	2 36 (82.8%) 49 (17.2%)
Stage of labour Latent Active Second	154 (54%) 87 (30.5%) 44 (15.4%)	Type of anesthesia Regional General	273 (95.8%) 12 (4.2)

Table	2	Obstetrical	Characteristics	of	Women	Undergoing	Emergency	Cesarean
Deliver	y a	at Selected H	lospitals of Nor	thw	est Ethio	pia, 2022/23 ((n=285)	

Abbreviations: ANC, Antenatal care; BCRP, Birth preparedness complication readiness plan; OR, Operation room.



Figure 2 Decision to delivery time among women underwent emergency cesarean delivery with different indications at selected hospitals of Northwest Ethiopia, 2023.



Rank of surgeon

Figure 3 Decision to delivery time among women who underwent emergency with the rank of surgeon at selected hospitals of Northwest Ethiopia, 2023.



Figure 4 The overall restricted mean decision to delivery time among women who underwent emergency cesarean delivery at selected hospitals of Northwest Ethiopia, 2023.

was shorter than those who delayed to accept at 54.9 min (95% CI: 52.7–57). With a p-value of 0.020, this difference was statistically significant (Figure 5).

The mean decision to delivery time for mothers with indication for cord prolapse is 32 min (95% CI: 28.7–35.3), which is shorter than with their courtparts such APH (39 min), obstructed labor (47 min), GIII MSAF (46.9 min), NRFHRP (48.8 min), CPD (50.4 min), two C/S scars (48.6 min), and others (51.4 min). This difference was statistically significant with p-value=0.000 (Figure 6).



Figure 5 Kaplan-Meier survival curve for comparison of the mean decision to delivery time among women who underwent emergency cesarean delivery with informed consent at selected hospitals of Northwest Ethiopia, 2023.



Figure 6 Kaplan-Meier survival curve for comparison of the mean decision to delivery time among women who underwent emergency cesarean delivery with indication at selected hospitals of Northwest Ethiopia, 2023.

The mean decision to the delivery time for women whose OR material was prepared was 48 min (95% CI: 46.3–49.8) which is shorter than their counterparts at 54.5 min (95% CI: 52.4–56.5), and testing equality among the groups with p-value of 0.006 was statistically significant among the group (Figure 7).

The mean decision to the delivery time for women whose free functional OR tables were present was 47.7 min (95% CI: 45.9–49.6) which is shorter with their counterparts at 54.3 min (95% CI: 2.5–56.2), and testing equality among the groups with p-value of 0.002 was statistically significant among the group (Figure 8).



Figure 7 Kaplan–Meier survival curve for comparison of the mean decision to delivery time among women who underwent emergency cesarean delivery with OR material prepared at selected hospitals of Northwest Ethiopia, 2023.



Figure 8 Kaplan-Meier survival curve for comparison of the mean decision to delivery time among women who underwent emergency with free functional OR table present at selected hospitals of Northwest Ethiopia, 2023.

The mean decision to the delivery time for women with general anesthesia was 37 min (95% CI: 30.2–44.3) which is shorter than for women with regional anesthesia at 49.5 min (95% CI: 47.9–51.1), and testing equality among the groups with p-value of 0.000 was statistically significant among the group (Figure 9).

Testing Proportional Hazards Assumption

A Cox regression model was used to examine the effects of sociodemographic, obstetrics, and treatment characteristics of patients on the decision to delivery time. The following variables were included in the model as predictors: residence, birth preparedness and complication readiness, consent, indication, bladder flap, working time, rank of the surgeon, types of anesthesia, OR material prepared, and free functional OR table present were all significantly associated with decision to delivery time. A goodness-of-fit (GOF) test was conducted to assess the proportional hazard (PH) assumptions of the



Figure 9 Kaplan-Meier survival curve for comparison of the mean decision to delivery time among women who underwent emergency cesarean delivery with regional anesthesia and general anesthesia at selected hospitals of northwest Ethiopia, 2023.

Cox model for given predictor variables. The findings indicated that all variables included in the model satisfied PH assumptions (p-value>0.05) with a global test of 0.5005 (Table 3).

After preserving the Schoenfeld and scaling the Schoenfeld, the proportional hazards assumption test residuals suggested that both covariate-specific and global tests are not violated (fulfilled). Hence, the Cox regression model is best for these data.

Predictors of Decision to Delivery Time

Following the testing of each variable in turn, the bivariable analyses revealed that residence, birth preparedness and complication readiness, consent, indication, bladder flap, working time, rank of the surgeon, types of anesthesia, time of transfer to OR, time of anesthesia, OR material prepared, and free functional OR table present were all significantly associated with the decision to delivery time.

After controlling confounders in multiple Cox regression analysis, consent, indication for cord prolapses, anesthesia types, free functional OR table present, rank of the surgeon, and bladder flap were independent predictors of the decision to delivery time. The multiple Cox regression analysis revealed that holding other variables constant, women who hesitate to accept informed consent are 83% less likely to have a decision to delivery time within 30 min (AHR: 0.17, 95% CI: 0.02–1.25) whenever there is a follow-up. Holding other variables constant, women with cord prolapse have 1.36 higher hazard of developing the decision to delivery time than those with other indications (AHR: 1.36, 95% CI: 0.46–3.94). Regarding the rank of surgeon, MSc in clinical midwifery students were 58% times less likely to decide on emergency C/S as compared to seniors (AHR: 0.42, 95% CI: 0.420.16–1.08). Moreover, relative to patients who have free functional OR tables present, those who have no free OR table were 73% times less likely to develop the decision to delivery time (AHR: 0.27, 95% CI: 0.28–0.94).

Furthermore, using a bladder flap during emergency C/S is 67% less likely to undergo the decision to delivery time as compared to not using (AHR: 0.33, 95% CI: 0.16–0.85). Emergency cesarean delivery with regional anesthesia is 44% less likely to have decision delivery time as compared with general anesthesia (AHR: 0.56, 95% CI: 0.25–1.28) (Table 4).

Discussion

In this cohort study, the aim is to determine the decision to delivery time and its predictors among women who underwent emergency cesarean delivery. At the end of three months of follow-up, 62 (21.8%) women delivered within 30 min or less with a mean DDT of 48.9 min, which is nearly similar with other studies of 19.6%, 21.3% in Gondar Southern Ethiopia, and 20.3% in Bahir Dar.^{14–16} This may be due to the similarity in the accessibility of logistics in hospitals, the practice and

Predictors	Rho	Chi-squared	df	Prob>chi-squared
Residence	-0.05543	0.61	I	0.4339
Working time	-0.05452	0.45	I	0.5042
Consent	0.10780	1.60	I	0.2054
OR material	0.15116	2.86	I	0.0908
Free functional OR table	-0.21817	6.94	I	0.084
Types of anesthesia	-0.05565	0.55	I	0.4573
Time of transfer	-0.05946	0.20	I	0.6513
Rank of surgeon	0.08596	0.47	I	0.4943
Time of anesthesia	-0.09540	0.62	I	0.4319
Global test		14.3	9	0.5005

 Table 3 Goodness-of-Fit Test Assessing Proportional Hazards Assumption

Variables	Survival Status		CHR (95% CI)	AHR (95% CI)	
	Delivered, N (%) Censored, N (%)				
Working time Day Night	44 (21.9%) 18 (21.4%)	157 (78.1%) 66 (78.3%)	। 0.99 (0.57–1.71)	I I.6 (I.04–2.47)	
Informed consent Agreed Hesitate	61 (23.6%) I (3.7%)	197 (76.4%) 26 (96.3%)	I 0.14 (0.02–1.02)	I 0.17 (0.02–1.25) **	
Indication for C/S APH	8(47.1%)	9(52.9%)	I	I	
Cord prolapses	12 (92.3)	l (7.7%)	1.65 (0.68-4.05)	1.36 (0.46–3.94)**	
Obstructed labor	2(18.2%)	9(81.8)	0.420 (0.09–1.98)	0.43 (0.084–2.15)	
GIII MSAF	6(16.2%)	31 (83.8%)	0.27 (0.93–0.77)	0.46 (0.14–1.56)	
NRFHR	21 (20.4%)	82 (79.6%)	0.35 (0.16-0.80)	0.66 (0.27–1.64)	
CPD Two C/D	4(12.9%) 2(68.9%)	27 (87.1%) 27 (93.1%)	0.19 (0.056–0.62) 0.09 (0.02–0.46)	0.29 (0.08–1.01)	
Others	7(44%)	37 (84.1%)	0.024 (0.09–0.66)	0.13 (0.03–0.64)	
Residence Urban Rural	46 (23.7%) 16 (17.6%)	148 (76.3%) 75 (82.4%)	l 0.71 (0.40–1.25)	l 1.34 (0.81–2.21)	
Rank of surgeon IESO MSc Clinical MW MSc in clinical MW students Senior	27 (31%) 12 (26.7%) 7(8.6%) 16 (22.2%)	60 (69%) 33 (73.3%) 74 (91.4%) 56 (77.8)	0.69 (0.37–1.29) 0.79 (0.40–1.56) 0.25 (0.109–0.58 I	1.49 (0.77–0.87) 2.88)0.87–0.91) 0.42 (0.161–1.21)** 1	
OR material Yes No	60 (24.3%) 2(5.3%)	187 (75.7%) 36 (94.7%)	l 0.18 (0.04–0.73)	l 0.33 (0.08–1.48)	
Free functional OR Yes No	59 (25%) 3(6.1%)	177 (75%) 46 (93.9%)	l 0.19 (0.06–0.62)	l 0.27 (0.28–0.94)**	
Types of anesthesia General Regional	55 (20.1%) 7(5.3% 55 (20.1%)	5(41.7%) 218 (79.9%)	l 0.27 (0.12–0.60)	I 0.56 (0.25–1.28)**	
Bladder flap Yes No	7(8.4.%) 55 (27.2%)	76 (91.6%) 147 (72.8%)	0.28 (0.12–0.62) I	0.33 (0.16–0.85)* I	
Time transfer ≤10 min >10 min	41 (27.7%) 21 (15.3%)	107 (72.3%) 116 (84.7%)	l 0.47 (0.28–0.79)	l 0.75 (0.42–1.34)	

Table 4 Cox Regression Analysis of Predictors of Decision to Delivery Time Among Women WhoUnderwent Emergency Cesarean Delivery at Selected Hospitals of Northwest Ethiopia, 2023 (N=285)

Notes: *Statistically significant at P value 0.25. **Significant at p value 0.05.

Abbreviations: Cesarean section, APH, Antepartum hemorrhage; GIII MSAF, Grade three meconium-stained amniotic fluid; CPD, Cephalopelvic disproportion; IESO, Integrated emergency surgeon officers; NRFHR, non reassuring fetal heart rate; MW, Midwifery.

experience of professionals, and types of cesarean delivery. The finding of this study is lower than in Saudi Arabia which was 30.6%, 87.5% in Denmark, 42.4% in India, 45% in Gondar in 2022, and in one study 50.29%, but is comparable with a study in Oman Qatar at 23.8%.^{17–22} This difference might be occurring due to various studies conducted in different with different set up of institutions and different sample sizes. The findings of this research were greater than those of studies conducted in Tanzania, Kenya, Benin teaching hospital, Nigeria, and Bahir Dar in 2021 which were 14.3%, 12%, 3%, 2.1%, and 5.7%, respectively.^{1,5,23–25} The difference might be due to general infrastructure, economic differences, multidisciplinary operative room team training, easy accessibility of logistics, and laboratory services.

In this study, the overall incidence of decision to delivery time was found to be 1.2 per 100-women minutes of observation. The mean decision to delivery time was 48.9 min (IQR=29–35; 95% CI: 52.5–61.7). This finding is higher than the studies done in Sunanda¹⁷, in Gondar¹⁶ by 2020, in Israel²⁶ and in Ireland,²⁷ in Bahir Dar,²⁵ and in Nigeria,⁵ in 2021. The difference might be the sample size, period, types of emergency cesarean delivery, and institutional differences. On the contrary, this finding is less than the study done in Benin teaching hospital¹ which was 106.3±79.5 min, in Indonesia,²⁸ in India,⁶ in Gondar,¹⁶ in Nigeria¹⁰ which was 145.3 ± 69.2 min, and in Norway²⁹ which was 52.4 min. The difference might be due to general infrastructure, economic differences, multidisciplinary operative room team training, easy accessibility of logistics, laboratory services, sample size, study period, and types of emergency C/S which include both categories in this study. So, an institution having an adequate operation table, experienced physician, anesthetist, operation room materials, operation room nurse, and urgency of indication could shorten the decision to delivery time interval in the emergency cesarean section. In this study, the presence of a free operation table, consent, indication for cord prolapses, rank of surgeon, anesthesia types, and use of a bladder flap are predictors of decision to delivery time in the multiple Cox regression analysis.

In this study, women who accepted informed consent had a statistically significant association with recommended DDT compared with women who hesitated to accept earlier. In this study, women who hesitated to accept informed consent were 87% less likely to have a decision to delivery time when compared to those who agreed to accept emergency C/S within 30 min. This finding is supported by a study done in Gondar.²¹ This study found that women whose EmCS was performed under regional anesthesia were 44% less likely to have the recommended DDT than women who were performed under general anesthesia. The findings of this study were in line with those of studies conducted in Israel,¹⁹ Indonesia, Norway, and Bahir Dar which showed that general anesthesia as a result of technical problems in inducing, and stabilization of clients before regional anesthesia is needed. Women whose emergency cesarean delivery was performed by seniors were more likely to have the recommended DDT than those made by MSc Clinical Midwife and MSc in Clinical Midwifery students. The same result was observed in Norway²⁹ and Bahir Dar.¹⁵ This could be because seniors were more experienced and had more exposure than others. A study in Singapore and Hong Kong showed no significant differences in DDI between senior surgeons and residents.

In this study, compared to women with a free OR table, those with no free functional OR table present were 73% less likely to have the recommended decision to delivery time. This finding is in line with studies conducted in different parts like Gondar,²¹ Gamo Zone,¹⁴ and Oman that revealed that women who had no additional OR table while they are preparing for EmCS were associated with a longer decision to delivery time. This might be attributed to the fact that if there is no additional OR table, it causes a prolonged decision to delivery interval secondary to the third delay. Regarding indication for emergency C/S, women with cord prolapse are 1.36 times more likely to have a decision to delivery time when compared with other indications. This may be due to the cord prolapses needing a prompt response from providers to save the life of the fetus. In this study around 92.3% of the study participants with cord prolapse had emergency C/S with recommended time. Another significant variable in this study is the use of a bladder flap which is 67% less likely to have the recommended delivery time within 30 min. This finding was supported by a study done in the USA which shortened the incision to delivery time among non-users.³⁰

Conclusion

The decision to delivery time among women underwent emergency cesarean delivery at DCSH and Lumame Primary Hospital was longer than the recommended interval of time. Only 21.8% of women were delivered within the

recommended DDT below 30 min. The presence of a free OR table, hesitated to consent, indication for cord prolapses, rank of surgeon, anesthesia types, and use of a bladder flap are predictors of decision to delivery time in the multiple Cox regression analysis. Despite the importance of doing EmCS as the recommended interval for the improvements of a perinatal outcome as well as quality care for mother and newborn, a substantial number of women had not achieved the recommended DDT (<30 min).

We strongly recommend that material which providers and facilities should be better equipped with a free OR table in advance and ready for a rapid emergency cesarean delivery.

Consent processes are also another important factor that determines the decision to the delivery time at Debre Markos Comprehensive Specialized Hospital and Lumame Primary Hospital. The care provider should give information at antenatal care about the possibility of operational delivery that helps to shorten time spent on the consent process. It would have been better if general anesthesia was used rather than regional anesthesia as a result of technical problems in inducing and stabilization of clients before regional anesthesia is needed. Also, it as better if ECD was performed by seniors because seniors were more experienced and had more exposure than others.

Abbreviations

ACOG, American College of Obstetrician and Gynecologist; ANC, Antenatal care; AHR, Adjusted hazard ratio; BPRC, Birth preparedness and complication readiness; CPD, Cephalo-pelvic disproportion; DCSH, Debre Markos Comprehensive Specialized Hospital; DDT, Decision to delivery time; CD, Cesarean delivery; EmCD, Emergency cesarean delivery; NRFHR, Non-reassuring fetal heart rate; OR, operation room; RCOG, Royal College of Obstetricians and Gynecologists.

Data Sharing Statement

Data will be available from the corresponding author on reasonable request.

Ethical Approval and Informed Consent

All methods were conducted according to the ethical standards of the Declaration of Helsinki. Ethical clearance was obtained from an institutional Research Ethics Review Board (IRB) of University of Gondar with reference number MIDW/30/2015 E.C. In addition, a letter of ethical approval was sent to Debre Markos Comprehensive Specialized Hospital and Lumame Primary Hospital and permission was obtained from the hospital's administrators. Informed, voluntary, written, and signed consent was obtained from the head of the hospital and mothers. There are no study participants under the age of 18 years. The standard safety measures for the prevention of COVID-19 were strictly followed throughout the data collection period. Before conducting the interviews, information was given to the participants and they were assured for voluntary participation, confidentiality, and freedom to withdraw from the study at any time. The nature and significance of the study was explained to the participants.

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Author Contributions

All authors made a significant contribution to the work reported, that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising, or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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Disclosure

The authors report no conflicts of interest in this work.

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