Perinatal outcomes of babies delivered by second-stage Caesarean section versus vacuum extraction in a resource-poor setting, Nigeria – a retrospective analysis

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Abstract

Background: To evaluate the perinatal status of neonates delivered by assisted vaginal delivery (AVD) versus second-stage caesarean birth (CS).

Methods: A 5-year retrospective study was conducted in a tertiary hospital. Data was analyzed with IBM SPSS® version 25.0 statistical software using descriptive/inferential statistics.

Results: A total of 559 births met the inclusion criteria; AVD (211; 37.7%) and second-stage CS (348; 62.3%). Over 80% of the women were aged 20–34 years: 185 (87.7%) for the AVD group, and 301 (86.5%) for the second-stage CS group. More than half of the women were parous: 106 (50.2%) for the AVD group, and 184 (52.9%) for the second-stage CS group. The commonest indication for intervention in both groups is delayed second stage: 178 (84.4%) in the AVD group, and 239 (68.9%) in the second-stage CS group. There was a statistically significant difference in decision to delivery interval (DDI) between both groups: 197 (93.4%) women in the AVD group had DDI of less than 30 min and 21 women (6.0%) in the CS group had a DDI of less than 30 min (p < 0.001). During the DDI, there were 3 (1.4%) intra-uterine foetal deaths (IUFD) in the AVD and 19 (5.5%) in the CS group (p = 0.023). After adjusting for co-variates, there were statistically significant differences between the AVD and CS groups in the foetal death during DDI (p = 0.029) and perinatal deaths (p = 0.040); but no statistically significant differences in severe perinatal outcomes (p = 0.811), APGAR scores at 5th minutes (p = 0.355), and admission into the NICU (p = 0.946). After adjusting for co-variates, use of AVD was significantly associated with the level of experience of the care provider, with resident (junior) doctors less likely to opt for AVD than CS (aOR = 0.45, 95% CI: 0.29–0.70).

Conclusion: Second-stage CS when compared with AVD was not associated with improved perinatal outcomes. AVD is a practical option for reducing the rising Caesarean delivery rates without compromising the clinical status of the newborn.

Keywords: Assisted vaginal delivery, Vacuum extraction, Neonatal assessment, Caesarean section, Decision to delivery interval, Second stage of labour
Background
Assisted vaginal delivery (AVD; vacuum extraction) and caesarean section (CS) are both obstetric procedures associated with enormous benefits and some complications to women and newborn babies. Either intervention can be undertaken during the second stage of labour for maternal and fetal indications, ranging from prolonged second stage, fetal distress, maternal exhaustion or maternal medical conditions [1, 2]. The procedure of choice should be individualized and depend on meeting pre-requisite criteria, decision-delivery-interval (DDI) and medico-legal considerations [1].

While caesarean section rates have increased dramatically worldwide in the last decades [3–6], AVD is significantly under-utilized, particularly in sub-Saharan Africa; accounting for a meager 1% of institutional births [7–10]. Indeed, the decreasing trend in the utilization of AVD in low and middle-income countries has largely been attributed to decreasing experience with skills required for AVD [7, 8]. Unnecessary CSs place an unjustified burden on the scarce financial and human resources that barely meet the health needs of low-income countries. Fortunately, there is clear evidence indicating that high caesarean section rates can be offset by using AVD, especially in resource poor settings where aversion for CS is paramount. Costs of CS are exorbitant and non-operational universal health insurance schemes results in catastrophic out-of-pocket payment for service charges [2, 7, 9, 11, 12].

Previously evaluated neonatal outcome of AVD was compared with baseline risk of complications from spontaneous vaginal delivery [1]. In situations, however, with life threatening maternal or fetal emergencies in second stage of labour, spontaneous vaginal delivery may not be the best comparison. Hence the need to compare the immediate newborn status, delivered by either AVD or CS. A prospective cohort study conducted in Uganda demonstrated that births by AVD had better maternal and equivalent perinatal outcomes compared to births by second-stage CS [10]. Given that there has been no similar study conducted in Nigeria or in the wider west African region, the present study was necessary to bridge this gap and provides non-existent local data. This study will hopefully complement already existing quality evidence and help improve decision making in obstetric units in resource-poor settings.

Methods
Study design
This was a five-year retrospective study, in which data were extracted from hospital records of women who had AVD and second-stage CS over a 5-year period (01 January 2012 to 31 December 2016) in the Federal Teaching Hospital of Abakaliki (FETHA).

Participants
Inclusion criteria were women with singleton pregnancy ≥37 weeks gestation, cephalic presentation, stage two of labour, engaged fetal head at station +2 and +3, and delivery assisted by vacuum device (Kiwi device®) or second-stage CS. Exclusion criteria were women with twin gestation, non-cephalic presentation, preterm births and women with uterine rupture before the decision to intervene. Women with fresh meconium-stained liquor with cardiotocographic evidence of fetal distress (deceleration, bradycardia, tachycardia or loss of baseline variability) in the first stage of labour were excluded.

Data collection
Women who had CSD were identified from the operating theatre book, and then their case notes were cross-checked to identify women who had fully dilated cervix before the decision to perform CS was made. Women who had AVD were identified from the delivery room register. Women who had CS due to failed AVD were included as AVD as this was the intention a priori. Data were extracted from the women’s case notes, and the admission and discharge register by three trained junior doctors using a proforma. Missing data were cross-checked with the corresponding nursing shift reports for the day. Socio-demographic data were extracted from the case notes. Other information extracted were Apgar scores (1st minute and 5th minute), adverse events during the interventions, admission into NICU and duration of admission. Indications for AVD and second-stage CS were classified as defined in Table 1. Decision to delivery intervals (DDI) were estimated from partographs, surgical notes, and nursing shift reports. Low Apgar score was defined as APGAR < 7 [13]. Women who experienced intra-uterine fetal death before the decision for intervention was made were excluded from the analysis.
of perinatal outcomes. Study outcomes were foetal death within the DDI, severe perinatal outcomes, low Apgar score at 5-min, admission into NICU, and perinatal death. Severe perinatal outcome was defined as the presence of any of the following: perinatal death, severe birth injury, 5-min Apgar score < 4, and/or convulsions. Severe birth injury was defined as presence of any of the following: dislocation of the leg, clavicular fracture, intracerebral haemorrhage and/or subgaleal haemorrhage.

Table 1 Definitions of indications for second stage intervention

| Indication classified as: | If the indication for the intervention as stated in the patients’ medical record is: |
|--------------------------|-------------------------------------------------------------------------------------|
| Delayed / Prolonged second stage (second stage of labour > 2 h) | Prolonged labour, obstructed labour, malposition, borderline or contracted pelvis or inadequate contractions |
| Foetal distress | Foetal distress, abnormal foetal heart rate, foetal tachycardia > 160/min or foetal bradycardia < 100/min |
| Maternal exhaustion / fatigue | Maternal exhaustion or insufficient maternal effort |
| Other indications | Other indication stated |

Statistical analysis

Data were entered into Microsoft Excel (Microsoft, Redmond, WA, USA), cleaned and transferred to SPSS version 25.0 (IBM, Armonk, NY, USA) for statistical analyses. Bivariate analyses were performed to determine associations between baseline maternal and neonatal characteristics and mode of delivery. When both Chi-Square test and Fischer’s Exact Tests were estimated, only the \( p \)-value of the Fischer’s Exact test is reported. Multivariate logistic regression analyses were performed to estimate adjusted odds ratios (aORs) with 95% confidence interval (CI) for study outcomes, and to also estimate the effect of the doctors’ obstetric experience on the odds of opting for a mode of delivery. \( P < 0.05 \) was used to define statistical significance and all tests were two-tailed.

Results

Amongst the 9287 births during the study period, 559 (6.0%) women met the inclusion criteria for the study; 211 (2.3%) with term singleton pregnancies had AVD and 348 (3.7%) women with term singleton pregnancies...
had second-stage CS. Thirteen women who had CSD due to failed AVD were included in the AVD group (Fig. 1).

Over four-fifth of women in both groups were in the age range 20–34 years and almost half were nulliparous women (Table 2). Other baseline sociodemographic characteristics of the women are shown in Table 2. The commonest indication for both AVD and CS was delayed (or prolonged) second stage of labour: 178 (84.4%) in the AVD group and 239 women (68.7%) in the CS group ($p < 0.001$).

Perinatal outcomes between both groups are shown in Table 3. Compared to 197 (93.4%) women in the AVD group, only 21 women (6.0%) in the CS group had a DDI of less than 30 min (OR 0.01; 95% CI 0.00–0.01).

There were 3 (1.4%) intra-uterine foetal deaths (IUFD) during the DDI in the AVD and 19 (5.5%) in the CS group (OR 0.25; 95% CI 0.07–0.85). Perinatal deaths were 7 (3.3%) in the AVD and 27 (7.8%) in the CS (OR 0.41; 95% CI 0.17–0.95) while severe perinatal outcomes were 22 (10.4%) in the AVD group and 42 (12.1%) in the CS group (OR 0.85; 95% CI 0.49–1.47).

Table 4 shows the results of multivariate logistics regression analysis for the main study outcomes. After adjusting for parity, booking status, gestational age, indication for the intervention, presence of imminent delivery complications and birthweight, identified from univariate analysis as significant co-variates, statistically significant differences between the AVD and CS groups were detected in foetal deaths during DDI (aOR = 0.22; 95% CI 0.06–0.89) and perinatal deaths (aOR 0.41; 95% CI 0.17–0.96). After adjusting for significant co-variates in a multivariate logistics regression model, the use of AVD was significantly associated with the level of experience of the care provider, with resident doctors less likely to opt for AVD (aOR 0.45; 95% CI 0.29–1.47).

Discussion

The present study showed that for women who needed assistance in the second stage of labour, there was no significant difference in immediate neonatal outcomes and admission into the NICU for both AVD and CS. This re-affirmed the findings from previous studies which showed no significant difference in immediate neonatal outcomes for AVD and CS [5, 8, 9]. However, it was at variance with a similar study in Israel, where Shmueli et al. reported that CS yielded poorer neonatal outcome than AVD [14].

Incidence of 2.3% AVD in this study is similar to 2.0% in Maiduguri, Nigeria, but higher than 1.5% reported in Enugu, Nigeria and 0.54% reported in Bauchi, Nigeria [10, 15, 16]. Elsewhere in Africa, our incidence of AVD is slightly lower than the 3.1% reported in Kumasi Ghana and 2.8% reported in Kampala, Uganda; but

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**Table 2** Baseline characteristics of study cohorts

| Characteristics                        | Assisted Vaginal Delivery (n = 211) | Second-stage Caesarean Delivery (n = 348) | P-value |
|----------------------------------------|------------------------------------|------------------------------------------|---------|
| Mothers’ age                           |                                    |                                          |         |
| < 20 years                             | 12 (5.7%)                          | 21 (6.0%)                                | 0.916   |
| 20–34 years                            | 185 (87.7%)                        | 301 (86.5%)                              |         |
| ≥ 35 years                             | 14 (6.6%)                          | 26 (7.5%)                                |         |
| Marital Status                         |                                    |                                          |         |
| Single                                 | 2 (0.9%)                           | 3 (0.9%)                                 | 1.000   |
| Married                                | 209 (99.1%)                        | 345 (99.1%)                              |         |
| Educational qualification              |                                    |                                          |         |
| Primary education & below              | 52 (24.7%)                         | 95 (27.3%)                               | 0.552   |
| Secondary education & above            | 159 (75.3%)                        | 253 (72.7%)                              |         |
| Religion                               |                                    |                                          |         |
| Christian                              | 208 (98.6%)                        | 345 (99.1%)                              | 0.677   |
| Muslim, ATR & others                   | 3 (1.4%)                           | 3 (0.9%)                                 |         |
| Prenatal care received                 |                                    |                                          |         |
| ≥ 4 ANC visits                         | 143 (67.8%)                        | 231 (66.4%)                              | 0.942   |
| 1–3 ANC visits                         | 42 (19.9%)                         | 73 (21.0%)                               |         |
| 0 ANC visit                            | 26 (12.3%)                         | 44 (12.6%)                               |         |
| Parity                                 |                                    |                                          |         |
| Nulliparous                            | 105 (49.8%)                        | 164 (47.1%)                              | 0.600   |
| Parous                                 | 106 (50.2%)                        | 184 (52.9%)                              |         |
| Gestational age at birth               |                                    |                                          |         |
| 37–39 weeks                            | 134 (63.5%)                        | 207 (59.5%)                              | 0.372   |
| ≥ 40 weeks                             | 77 (36.5%)                         | 141 (40.5%)                              |         |
| Sex of newborn                         |                                    |                                          |         |
| Female                                 | 107 (50.7%)                        | 169 (48.6%)                              | 0.663   |
| Male                                   | 104 (49.3%)                        | 179 (51.4%)                              |         |
| Birth weight                           |                                    |                                          |         |
| < 2500 g                               | 13 (6.3%)                          | 14 (4.3%)                                | 0.561   |
| 2500 to 4000 g                         | 178 (85.5%)                        | 285 (86.6%)                              |         |
| ≥ 4000 g                               | 17 (8.2%)                          | 30 (9.1%)                                |         |
| Indications for intervention a         |                                    |                                          |         |
| Delayed second stage                   | 178 (84.4%)                        | 239 (68.7%)                              | < 0.001 |
| Maternal exhaustion / fatigue          | 49 (23.2%)                         | 63 (18.1%)                               | 0.157   |
| Foetal distress                        | 6 (2.8%)                           | 71 (20.4%)                               | < 0.001 |
| Other indications                      | 0 (0.0%)                           | 8 (2.3%)                                 | 0.027   |
| Imminent delivery complications        |                                    |                                          |         |
| Impending Uterine rupture              | 3 (1.4%)                           | 9 (2.6%)                                 | 0.393   |
| Placenta abruption                     | 1 (0.5%)                           | 1 (0.3%)                                 | 1.000   |
| Cord prolapse                          | 1 (0.5%)                           | 7 (2.0%)                                 | 0.166   |

**Abbreviations:** ATR African traditional religion, ANC Antenatal Care consultations

*a* More than one indication could apply
Severe Birth injury includes any of the following: dislocation of the leg, clavicular fracture, intracerebral haemorrhage or subgaleal haemorrhage. More than one adverse event could apply in the first week after delivery.

Severe Perinatal Outcome defined as presence of the following: perinatal death, severe birth injury, 5-min Apgar score < 4, or convulsions. Only surviving newborns included for this analysis (Second-stage Caesarean delivery, n = 208).

Table 3 Delivery Outcomes & Perinatal outcomes of Surviving newborns

| Decision to Delivery Interval | Assisted Vaginal Delivery (n = 211) | Second-stage Caesarean Delivery (n = 348) | Crude OR (95% CI) |
|------------------------------|-------------------------------------|------------------------------------------|-----------------|
| < 30 min                     | 197 (93.4%)                         | 21 (6.0%)                                | 0.01 (0.00–0.01) |
| ≥ 30 min                     | 14 (6.6%)                           | 327 (94.0%)                              |                 |

Outcome of delivery

- IUFD during DDI 3 (1.4%) 19 (5.5%) 0.25 (0.07–0.85)
- Perinatal death 7 (3.3%) 27 (7.8%) 0.41 (0.17–0.95)
- Severe Perinatal outcome 22 (10.4%) 42 (12.1%) 0.85 (0.49–1.47)

Timing of death

- During DDI 3 (1.4%) 19 (5.5%) 0.25 (0.07–0.85)
- Early Neonatal period 4 (1.9%) 8 (2.3%) 0.82 (0.24–2.76)

Apgar scores at 5 min

- ≥ 7 184 (88.5%) 294 (89.4%) 1.10 (0.63–1.90)
- 4–6 4 (2.4%) 29 (8.8%) 0.78 (0.46–1.33)
- < 4 3 (1.4%) 6 (1.8%) 0.85 (0.49–1.47)

Admission into the NICU

- Total admissions 30 (14.4%) 46 (14.0%) 1.04 (0.63–1.70)
- Length of Stay, > 7 days 5 (2.4%) 7 (2.1%) 1.13 (0.36–3.62)

Adverse events

- Birth asphyxia 13 (6.3%) 16 (4.9%) 1.30 (0.61–2.77)
- Convulsions 5 (2.4%) 8 (2.4%) 0.99 (0.32–3.06)
- Jaundice 5 (2.4%) 6 (1.8%) 1.33 (0.40–4.40)
- Sepsis 6 (2.9%) 9 (2.7%) 1.06 (0.37–3.01)
- Breathing difficulties 3 (1.4%) 4 (1.2%) 1.19 (0.26–5.37)
- Feeding difficulties 1 (0.5%) 2 (0.6%) 0.79 (0.07–8.77)
- Severe Birth injury 3 (1.4%) 1 (0.3%) 4.80 (0.50–46.46)

Table 4 Multivariate logistics regression analysis of perinatal outcomes

| Perinatal outcomes | Second-stage Caesarean delivery Adjusted OR (95% CI) | Assisted Vaginal delivery Adjusted OR (95% CI) |
|--------------------|------------------------------------------------------|-----------------------------------------------|
| Fetal Death within DDI | – Adjusted OR ² 1.00 (Reference) 0.22 (0.06–0.89) |                                 |
| Perinatal death | – Adjusted OR ² 1.00 (Reference) 0.41 (0.17–0.96) |                                 |
| Severe Perinatal Outcomes | – Adjusted OR ² 1.00 (Reference) 1.07 (0.60–1.92) |                                 |
| Low APGAR score at 5 min | – Adjusted OR ² 1.00 (Reference) 0.78 (0.46–1.33) |                                 |
| Admission into NICU | – Adjusted OR ² 1.00 (Reference) 0.98 (0.58–1.66) |                                 |

Abbreviations: OR Odds ratio, CI Confidence interval, IUFD Intrauterine Foetal Death, DDI Decision to delivery interval, NICU Neonatal Intensive Care Unit

Higher than the overall average of about 2.0% reported in rural Tanzania [2, 8, 11, 17]. On the other hand, our incidence of second-stage CS (3.7%) is higher than the 3.3% reported in Kampala, Uganda [10].

The need for interventions to resolve feto-maternal complications in the second stage of labour is necessary. The current study showed that prolonged second stage of labour was the most common indication for 84.4% AVDs and 68.7% CSs. Other researchers have also reported similar indication for AVD and CS [2, 10, 18, 19]. However, other studies have also reported poor maternal effort/exhaustion and fetal distress as major indications for second stage interventions. Opoku et al. reported an incidence of 10.9% for maternal exhaustion and 15.4% for fetal distress in AVD [1, 2].

Several studies have reported low incidence of AVD in many countries, including low-income countries of sub-Saharan Africa [1, 3, 9, 10, 12]. However, the recent increase in Caesarean section rates in many countries has not resulted in any significant improvement in neonatal outcomes [20]. In addition, several high-quality studies demonstrate that the use of AVD in well-selected patients remains a safe and effective method of delivering healthy neonates without compromising the overall birthing experience and outcome [8, 11, 21]. AVD is therefore a safe alternative worldwide, but more so in low-resource settings, where there is high aversion to CS due to several socio-cultural reasons [1, 10, 15, 22].
therefore disturbing that even for similar indications and similar set of women, resident doctors were less likely to opt for AVD than CS. This is mostly due to decreasing residents doctors’ experience in AVD [1, 10]. If this low usage of AVD persists, it may further aggravate the increasing rates of CS that are already very high in this setting [22]. Hence, given that mothers in this setting have very high aversion to CS and mothers who had AVD report better overall quality of life than mothers who had CS [15, 21], it is imperative that AVD should be encouraged where conditions allow.

Although this study significantly adds to the body of evidence comprehensively contrasting perinatal outcomes for newborns born via AVD with newborns born by second-stage CS, it is not without some limitations. This study did not present similar analysis on maternal outcomes. Also, this study was also performed in one hospital, which could limit it applicability to other settings. There is, therefore, a need for a large multi-centre study with longer follow-up period, incorporating other methods of assessing immediate neonatal outcomes such as arterial blood gas, and qualitative studies to understand the factors driving resident (junior) doctors’ decision-making in second stage events and how these can be tackled to improve usage of AVD in these settings.

Conclusions
AVD compared with second-stage CS was not associated with worse perinatal morbidity and mortality. Junior doctors are short in confidence in the use of a vacuum device for AVD. With appropriate trainings, AVD could be a practical option in reducing the rising caesarean delivery rates without compromising the clinical status of the newborns.

Abbreviations
AVD: Assisted vaginal delivery; CS: Caesarean section; FETHA: Federal Teaching Hospital Abakaliki; NICU: Newborn Intensive Care Unit

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Authors’ contributions
PE and OLL designed the study, oversaw its conduct, data analysis, interpretation of results, drafted the original article and reviewed the final draft. RUC contributed to the design of the study, collection and analysis of the data, and review of the article. CIU assisted in the drafting of the article, data analysis, interpretation and review of the article. CAl participated in the design of the study, collection, analysis and interpretation of data as well as the review of the final manuscript. All the authors read and approved the final draft of the article.

Availability of data and materials
The data that support the findings of this study are available from Federal Teaching Hospital Abakaliki, but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors upon reasonable request and with permission of Federal Teaching Hospital Abakaliki.

Ethics approval and consent to participate
Ethical approval was granted by the Ethics review committee of Federal Teaching Hospital, Abakaliki (FETHA), who determined the study did not constitute human subject trial and granted ethical approval for the study (FETHA/REC/Vol1/2018016).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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