Timing of Antenatal Corticosteroid Administration in Monoamniotic Twins

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Abstract

Objective  This study was aimed to determine if different strategies of antenatal corticosteroid (ACS) administration in monoamniotic twins leads to receipt within 7 days of delivery.

Study Design  This is a retrospective cohort of monoamniotic twins managed at a single institution from 2007 to 2017. Patients were classified as to whether ACS were administered upon admission or at a predetermined gestational age (grouped together as “routine”) or for a change in clinical status (“indicated”). We used univariate analyses to associate ACS administration strategies with our primary outcome: receipt of ACS within 7 days of delivery. We then used generalized estimating equations to examine associations between fetal monitoring patterns and delivery within 1 week.

Results  Twenty-four patients were included: eighteen patients in the “routine” group and six patients in the “indicated” group. There was no difference in optimal timing of ACS administration. Women experiencing delivery within the week were thrice more likely to spend on average more than 3 hours/day on the fetal monitor when compared with those who remained undelivered.

Conclusion  Administration of ACS on admission is not effective. Fetal heart rate tracing surveillance might be a better methodology to predict delivery and guide ACS administration.

Keywords
► monoamniotic twins
► antenatal corticosteroids
► preterm delivery
► antenatal fetal testing

Monoamniotic twin pregnancies are rare, comprising 1% of all twins and 5% of monochorionic twins.1,4 Monoamniotic twins are uniquely associated with a significant risk of fetal death due to prolonged cord occlusion from cord entanglement as both twins share the same amniotic sac. This cord entanglement is almost inevitable and present in nearly all gestations.3,4 Literature, published more than 10 years ago, had reported a perinatal mortality rate as high as 30 to 70%.5,6 However, more contemporary systematic reviews that have excluded anomalous fetuses and complications from twin-to-twin transfusion syndrome suggest a lower rate of 10 to 12%.7,8 Prenatal diagnosis, closer fetal surveillance, and elective preterm delivery have been attributed to this lower mortality rate. Prior retrospective research supports that inpatient admission with frequent fetal testing is better at preventing intrauterine fetal death than outpatient management.5,9,10 However, there is still debate about the optimal antenatal management and delivery timing for these patients.

Given the significant risk of intrauterine death, elective premature delivery is guaranteed for all pregnancies with most data supporting cesarean delivery at 32 to 34 weeks.5,9,11 The administration of antenatal corticosteroids (ACS) between 23 and 34 weeks has shown to reduce the risk of neonatal death, respiratory distress syndrome, and neurological injury.12 Even though there is a paucity of research regarding the use of ACS in multiple pregnancies, administration is
recommended regardless of fetal number. The original trial of Liggins and Howie demonstrated the need for ACS to be given within 1 week of delivery to be clinically efficacious. These findings have been corroborated by several trials.

Optimal timing of ACS is particularly relevant in monoamniotic twins since their prematurity is inevitable. Prior studies have focused on neonatal outcomes with a wide variation on fetal testing strategies but the specific information regarding the rate of administration, protocol, and optimal timing of ACS is often lacking.

We were interested to determine if different strategies for antenatal corticosteroid (ACS) administration in monoamniotic twin pregnancies would lead to optimal timing of ACS administration prior to delivery. Our hypothesis was that administration of ACS upon admission to the hospital or at a predetermined gestational age (< 30 weeks) would not lead to receipt of steroids within 7 days of delivery. Furthermore, we wanted to demonstrate whether our cohort of inpatient monoamniotic twin management would elucidate any predictors for the timing of delivery.

Materials and Methods

This was a retrospective cohort study of monochorionic monoamniotic twin pregnancies at or beyond 24 weeks of gestational age that were managed at a single tertiary care center from 2007 to 2017. Potential cases of monoamniotic twin gestation were identified using hospital birth records. Inclusion criteria were a live monochorionic-monoamniotic twin gestation at time of admission, admission to the hospital between 24 and 28 weeks of gestation and delivery at our institution. Those patients who had a monoamniotic twin pregnancy due to iatrogenic or spontaneous intertwin membrane rupture were excluded. Conjoined twins were also excluded. The time of hospital admission was decided between the parents, Obstetric provider, and neonatologists. Parents chose a gestational age at which they would want neonatal resuscitation if delivery was indicated.

All patients were admitted to the antepartum floor of our hospital where they were managed by the antepartum team which consisted of resident, fellows and a maternal–fetal medicine (MFM) specialist. As per hospital protocol, the antepartum fetal monitoring consisted of either twice (2007–2012) or thrice (2013–2017) daily fetal heart rate monitoring. The fetal heart rate monitoring consisted of 1 hour of fetal heart tracing each time the fetal heart rate was monitored (either twice or thrice daily). Additional monitoring was performed due to a finding on routine testing. If there were repetitive variable or late decelerations or the fetal heart tracing was difficult to interpret (discontinuous), the patient had prolonged fetal monitoring (the number of hours depending on the indication or concern of the obstetric provider). Repetitive decelerations were defined as more than two in a 30-minute period. Ultrasounds were performed weekly and fetal growth was estimated every 2 weeks. The degree of cord entanglement on ultrasound was noted but it was not an indication for additional fetal monitoring or delivery. Doppler’s velocimetry studies were only performed routinely when there were growth abnormalities and not to screen for umbilical cord compression. There was no protocol regarding timing of steroid administration, this was provider dependent.

Our primary exposure of interest was the strategy for the first course of ACS administration. We classified patients as to whether they received ACS on admission or at predetermined gestational age (grouped together as “routine”), or if ACS were reserved until a change in maternal or fetal clinical status (“indicated”). The change in clinical status was documented in the medical record. Our primary outcome of interest was receipt of ACS within 7 days of delivery. We made note of both first and repeat courses of ACS. Additional secondary outcomes of interest included rates of neonatal morbidity based on the work by Barrett and colleagues. We analyzed composite neonatal morbidity for the pregnancy overall and separately for presenting and nonpresenting twins. The composite neonatal outcome included one or more of the following: intubation for 2 or more days, birth trauma (including cephalohematoma, clavicular fracture, or long-bone fracture), infection requiring antibiotics, intraventricular hemorrhage (IVH), hypoxic ischemic encephalopathy, and death. We made note of need for intubation though this was excluded from our composite neonatal outcome if the duration of intubation was less than 2 days.

The associations between ACS administration strategy and primary and secondary outcomes were analyzed with Chi-square test or Fisher’s exact test for categorical variables and Wilcoxon’s rank sum test for continuous variables. Based on the results of our univariate analysis and our small sample size multivariate analysis was deferred. We next performed a secondary analysis to examine clinical factors associated with delivery within 1 week. For this analysis, we considered each week of patient hospitalization including the monitoring needs of each patient. We recorded daily hours on the fetal heart rate monitor for each woman throughout her inpatient hospitalization. Our outcome of interest for this secondary analysis was delivery within the week of interest. For this analysis, we first used univariate analysis to look for associations between clinical characteristics of interest including fetal monitoring patterns and delivery within the week. We then used generalized estimating equations to generate odds ratios examining the associations between fetal monitoring patterns and delivery while accounting for correlations in data over time.

Demographic, obstetric, and labor and delivery data were collected by chart review. Study data were collected and managed using REDCap electronic data capture tools hosted by the institution. All analyses were performed with Statistical Analysis Software (SAS), Version 9.4 (Copyright 2013–2017, SAS Institute, Inc, Cary, NC). The study was approved by our Institutional Review Board (Protocol number 2016P002057, approved on 11/1/2016).

Results

During the study period of 2007 to 2017, 27 patients with a monochorionic-monoamniotic twin gestation delivered at our institution. Three patients were excluded; two of whom...
were transferred from outside hospitals for delivery and one as both twins had died at 25 weeks (prior to a planned admission at 26 weeks). - Fig. 1 shows the inclusion, classification, and analysis of patients. All patients in the cohort received a complete course of ACS. Eighteen patients received ACS upon admission (24–28 weeks) or at a predetermined gestational age (26–30 weeks), and six patients received ACS in the setting of a change in clinical status during the hospital admission. The change in clinical status \((n = 6)\) encompassed four cases of fetal-heart rate tracing documented as nonreassuring that required prolonged monitoring, one new diagnosis of intrauterine growth restriction, and one case of preterm premature rupture of membranes. Seventy-nine percent (19/24) of the patients received one complete repeat course of repeat ACS. The repeat course of ACS (only one course) was given due to nonreassuring fetal heart tracing \((n = 7)\), concern for preterm labor \((n = 1)\), or because the pregnancy was still undelivered after 30 weeks and there had been no recent ACS exposure \((n = 11)\). There were no differences in maternal/obstetrics characteristics between these groups (Table 1). There were also no differences in neonatal outcomes (Table 2).

The median gestational age at delivery of our cohort was 32.3 weeks (range: 28.1–35.3 weeks). All patients in the cohort were delivered by cesarean section. The indications for delivery were scheduled (eight), nonreassuring fetal heart tracing

**Table 1** Maternal and obstetric characteristics per antenatal corticosteroid administration strategy

| Characteristics                      | Routine ACS \((n = 18)\) | Indicated ACS \((n = 6)\) | \(p\)-Value |
|--------------------------------------|---------------------------|---------------------------|-------------|
| Maternal age (y)                     | 32 (28–38)                | 35 (31–41)                | 0.40        |
| Nulliparous                          | 9 (50.0)                  | 2 (33.3)                  | 0.64        |
| Spontaneous conception               | 14 (77.8)                 | 4 (66.7)                  | 0.62        |
| Intrauterine growth restriction on ultrasound | 5 (27.8)              | 2 (33.3)                  | 1           |
| Cord entanglement on ultrasound      | 18 (100.0)                | 6 (100.0)                 | 1           |
| TTTS                                 | 0                         | 0                         | 1           |
| Gestational age at admission (wk)    | 26.5 (25–28)              | 27.3 (26.3–27.7)          | 0.37        |
| Gestational age at delivery (wk)     | 32.3 (32.0–33.1)          | 32.5 (29.9–34)            | 0.95        |
| ACS                                  | 18 (100.0)                | 6 (100.0)                 | 1           |
| Second course of ACS                 | 16 (88.9)                 | 3 (50.0)                  | 0.08        |

Abbreviations: ACS, antenatal corticosteroids; TTTS, twin-to-twin transfusion syndrome.

Note: Data are shown as \(n\) (%) or median (interquartile range).
status (13), preterm labor (two), and preeclampsia (one). There were no stillbirths in this cohort. All patients had cord entanglement reported on their ultrasounds. There was one neonatal demise at 2 days of life due to severe multiple congenital anomalies. This twin was born at 31.6 weeks with multicystic dysplastic kidneys, developed severe renal failure, and the parents opted to withdraw care. The mortality rate of our cohort was 2.1% (95% confidence interval [CI]: 0.1–11.1%; 0% if anomalous fetuses were excluded). The other anomalies in the cohort were cardiac, renal, and pulmonary.

- Table 3 demonstrates that there was no difference in optimal timing of ACS administration (receipt within 7 days of delivery) regardless of the ACS administration strategy. None of the patients in either group received their first course of ACS within 1 week of delivery. There was no difference between the groups in the rate of second course of ACS within 7 days of delivery. Furthermore, even those patients who received the first course of ACS based on a clinical change in status did not deliver within 7 days. The rate of the second course of ACS administration within 7 days of delivery was higher in the routine ACS group (62.5%) than the indicated ACS group (50%); however, this difference was not significant. Our results showed that none of the strategies used in our hospital led to optimal administration of ACS. Upon chart review of our records, it became apparent

Table 2 Neonatal outcomes per antenatal corticosteroids administration strategy

| Outcome                  | Routine ACS (n = 36) | Indicated ACS (n = 12) | p-Value |
|--------------------------|----------------------|------------------------|---------|
| NICU admission           | 36 (100.0)           | 12 (100.0)             | 1       |
| Birth weight twin A (g)  | 1,757.5 (1,587–1,927)| 1,589 (1,250–2,055)    | 0.48    |
| Birth weight twin B (g)  | 1,743 (1,701–1,900)  | 1,666 (1,285–1,920)    | 0.5     |
| Composite twin A*        | 5 (27.8)             | 1 (16.7)               | 1       |
| Composite twin B*        | 5 (27.8)             | 1 (16.7)               | 1       |
| Stillbirth               | 0                    | 0                      | N/A     |
| Neonatal death           | 0                    | 1 (8.3)                | 0.25    |
| Fetal anomaly            | 2 (5.55)             | 3 (25)                 | 0.09    |

Abbreviations: ACS, antenatal corticosteroids; N/A, not available; NICU, neonatal intensive care unit.
*There were no cases of intraventricular hemorrhage, birth trauma, infection, hypoxic ischemic encephalopathy.
Note: Data are shown as n (%) or median (interquartile range).

Table 3 Administration of antenatal corticosteroids within 7 days of delivery per administration strategy

| Outcome                  | Routine ACS (n = 18) | Indicated ACS (n = 6) | p-Value |
|--------------------------|----------------------|-----------------------|---------|
| 1st course of ACS within 7 d of delivery | 0 | 0 | 1 |
| Repeat course of ACS     | 16 (88.9)            | 4 (66.7)              | 0.25    |
| Repeat course of ACS within 7 d of delivery | 10 (62.5) | 2 (33.3) | 1 |
| Any ACS within 7 d of delivery | 10 (55.6) | 2 (33.3) | 0.64 |

Abbreviation: ACS, antenatal corticosteroids.
Note: Data are shown as n (%) or median (interquartile range).
that the change in clinical status was often related to instances where prolonged fetal heart rate monitor occurred but not always. Our analysis showed that patients who had been on the fetal monitor on average more than 3 hours a day were thrice more likely to be delivered within that week. This objective measurement, rather than just a subjective clinical change, is a novel clinical tool to better time ACS administration prior to delivery.

Whether umbilical cord occlusion can be anticipated is debatable. Given this unpredictability, most specialists recommend ACS administration upon admission. However, this very unpredictability and the potential fetal mortality associated with placental insufficiency and placental vascular anastomoses support a clinically driven approach. Our study suggests that changes in fetal heart rate patterns are associated with delivery within that week. Specifically, those patients who spent on average more than 3 hours a day on the fetal heart rate monitor were thrice more likely to be delivered that week. Even though we cannot conclude that intrauterine fetal death can be prevented with fetal surveillance, our results suggest that routine intermittent fetal surveillance leading to prolonged monitoring in a day is a harbinger of delivery within that week, giving a logical rationale for ACS administration.

The perceived importance of receiving ACS before delivery, combined with fear of missing the opportunity altogether, may overshadow the actual need for the medication to be given within 1 week of delivery to be efficacious. In our cohort, there were only six patients (25%) who delivered prior to 32 weeks and all of them spent a prolonged time on fetal heart rate monitoring prior to delivery (median of 12.5 hours, range: 7.5–21.25 hours). From these results, one can assume that the need for prolonged fetal monitoring raises enough clinical concern that would allow for timely ACS administration prior to delivery. For those patients in the “indicated” ACS group, 50% delivered before 32 weeks of gestation and all three of them had a complete second course of ACS. Although small numbers, the fact that no patients have missed administration ACS is reassuring as to the appropriateness of incorporating the fetal heart rate tracing interpretation into clinical management of ACS administration. By using this objective change in fetal monitoring pattern, we could increase the rate of optimal ACS administration and potentially improve neonatal outcomes.

Table 4 Fetal heart rate monitoring characteristics according to delivery within the week

| Characteristic                          | Undelivered (n = 126) | Delivery within 7 days (n = 24) | OR (95% CI) |
|----------------------------------------|-----------------------|-------------------------------|------------|
| Percentage change in h on monitor      | 0 (–0.02 to 0.07)     | 0.06 (–0.03 to 0.70)          | 4.12 (1.67–10.14) |
| Increase in wkly h by 10%              | 29 (23.0)             | 12 (50.0)                     | 3.34 (1.56–7.19) |
| Median daily h of monitoring           | 3.0 (2.14–3.14)       | 3.2 (2.0–5.0)                 | 1.52 (1.19–1.94) |
| Median > 3 h of daily monitoring       | 35 (27.8)             | 13 (54.2)                     | 3.07 (1.37–6.87) |

Abbreviations: CI, confidence interval; OR, odds ratio.
Note: data are shown as n (%) or median (interquartile range).
Of patients admitted for antenatal surveillance, we had no cases of stillbirths and only one neonatal demise from complications of multiple congenital anomalies. We decided to exclude cases of pregnancies affected by a stillbirth prior to admission (n = 1) because our primary outcome was optimal timing of ACS administration and the focus of our analysis was not on neonatal morbidity or timing of delivery as in most published literature. The very low mortality rate of our institution (2.1 or 0% if anomalous fetuses were excluded) is consistent with prior research published by Prefumo et al and Baxi and Walsh.7,22 There is a wide variation in mortality rate reported in the literature with older data suggesting a mortality rate as high as 70% which is often attributed to the fact those studies included demises due to fetal anomalies and fetal losses that were prior to 20 weeks.5,6

One of the strengths of our study is that even though it is a small case series, we gathered individual patient data regarding fetal heart rate monitoring. Our results provide novel data regarding an objective change in fetal heart rate monitoring that is associated with delivery within that week.

Limitation

Our study is not without limitations. Its retrospective nature being the largest. Another limitation is our small sample size and thus, we are likely underpowered to detect a difference in ACS administration strategies. Using the outcome of any ACS within 7 days of delivery, with rates of 55.6% in the routine group and 33.3% in the indicated group, and an α of 5% a study would need 60 patients in each arm to detect a difference with 80% power. Given the rarity of monoamniotic twins and the underlying risk of stillbirth associated with these pregnancies, a randomized control trial of strategies for ACS administration would be extremely difficult to accomplish. Given the small sample size of our study, we cannot generalize our findings and a large-scale multicenter study is essential to corroborate our findings. Another limitation of our study is that our cohort of patients was established from hospital birth records. Historical records are never as accurate as prospective planned data collection.

Conclusion

In conclusion, our results demonstrate that ACS administration upon admission does not lead to optimal ACS administration. The use of inpatient fetal surveillance with the opportunity of prolonged fetal heart rate monitoring when clinically indicated might be a better methodology to predict delivery timing and likely better time ACS administration.

Note

The study was performed at Brigham and Women’s Hospital, Boston, MA.

Presentation

These findings were presented as a poster at the 38th Annual Scientific Meeting of the Society for Maternal–Fetal Medicine in Dallas, TX, February 2018.

Conflict of Interest

The authors report no conflicts of interest and there was no financial support provided for this project.

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