A programmed labour protocol for optimizing labour and delivery

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Received: 14 February 2015
Revised: 24 February 2015
Accepted: 01 March 2015

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ABSTRACT

Background: Objectives: 1) To evaluate the effect of programmed labor on duration of labor 2) To assess efficiency of analgesics in reducing severity of labor pains and 3) To find out any maternal and fetal/neonatal complications.

Methods: 100 primigravida pregnant women meeting inclusion criteria admitted in labor room 50 were randomly enrolled in study group and 50 in control group.

Results: Significant reduction in duration of active phase of labor and 43 (86%) cases had pain relief. Out of them 20 (40%) cases experienced excellent pain relief while in controls, only 22% had pain relief.

Conclusions: Labour analgesia ensures pain relief, avoid maternal hyperventilation controls alteration of placental circulation thereby safe guarding the fetus against hypoxia. Thus programmed labor is easy, simple and effective method for painless delivery

Keywords: Programmed labour, Analgesia, Pain relief

INTRODUCTION

Labour is defined as the progressive dilatation of uterine cervix with co-ordinate uterine contractions that effect in and expulsion of the products of conception. These uterine contractions are perceived as labor pains. Labour is physiological but painful event. The anxiety, fear and stress a women suffers is beyond description. Labour analgesia ensures pain relief, controls alteration of placental circulation thereby safe guarding the fetus against hypoxia.

Programmed labor concept: This concept rest on three pillars;

1. Providing optimum pain relief - Use of analgesics and antispasmodics.
2. Ensuring adequate uterine contractions - Active management of labor
3. Close clinical monitoring of labor events - Maintaining a PARTOGRAM

Currently, the proven obstetric analgesia is epidural anesthesia. In places where epidural analgesia cannot be provided, tramadol, a centrally acting non opioid analgesic has been used as a labor analgesic. Ketamine, a dissociative anesthetic, is gaining popularity as it provides excellent pain relief and patient satisfaction. The present study attempts to compare the duration of labor and maternal outcome in programmed labor protocol and conventional labor protocol followed in our hospital.

Aims and objectives

1) To evaluate the effect of programmed labor on duration of labor.
2) To assess efficiency of analgesics in reducing severity of labor pains.
3) To find out any maternal and fetal/neonatal complications.

**METHODS**

The present prospective randomized study was undertaken at national institute of medical sciences Jaipur (India). It was approved by ethical committee of the institute.

The patients in active labor were divided into two groups by simple randomization.

Group I: 50 (cases) for programmed labor.

Group II: 50 (controls) for traditional management of labor.

Total sample size was 100 primigravidas.

On admission to Labor room detailed history was taken and a thorough physical and general examination was done. Obstetrical examination including per vaginal examination was done and pelvic assessment was done to rule out cephalopelvic disproportion, after confirmation subject in active labor ARM was done for confirmation of colour of liquor, cases were selected for study. All subjects were subjected for routine investigation.

Every women were counseled regarding drugs used in programmed labor and after counseling written informed consent was taken.

Level of analgesia assessed using following scale:

0 - No pain relief
1 - Mild pain relief
2 - Moderate pain relief
3 - Excellent pain relief

**Inclusion criteria**

1. Age, between 21-35 years.

2. No identifiable medical or obstetric complications present.

3. Primigravida with singleton pregnancy with cephalic presentation with spontaneous onset of labor.


5. Admission NST-reactive.

6. Active phase of labor with cervical dilatation 4cms and 50% effaced.

7. Liquor should be clear after ARM.

8. No clinical evidence of cephalopelvic disproportion.

**Exclusion criteria**

1. High risk cases like antepartum haemorrhage, preeclampsia, diabetes complicating pregnancy, polyhydramnios, oligohydramnios, cephalopelvic disproportion, malpresentation, and pre labor rupture of membranes.

2. Patient who are not willing to sign informed consent will be excluded from the study.

**Protocol**

- The cervix should be 3.0-4.0 cm dilated, >50% effaced and head is at 0 or -1 station
- Amniotomy is performed at 3-4 cm dilatation.
- Start an intravenous infusion line with 5% Ringer Lactate solution @ about 20 drops/min
- Ensure that pains are optimal that is 3-4 c/35-45”/10’.
- If needed half an hour after amniotomy, Oxytocin drip 5units in 500ml RL started at 8-10drops / min and titrated every 30mins up to adequate contractions (3-4 c/ 35-45”/10’) achieved.
- Inj. tramadol 1 mg/kg and inj. drotaverine hydrochloride 40 mg I.M. single dose is given at amniotomy.
- 2 mg of diazepam + 6 mg pentazocine given i.v. at amniotomy and repeated 2 hourly on patient demand.
- Progress of labour monitored by partogram and p/v examination done every 2 hourly after amniotomy.
- 10 units of oxytocin diluted in 20 ml of saline and given through the umbilical vein of the placenta or administer it slow intravenous to the mother.
- Or
  - Inj. prostadin 125 mcg given IM after birth of the baby.
- Duration of active phase of labor, 2nd stage and 3rd stage of labor will be assessed.
- Neonatal assessment is done with APGAR score at 1min and 5 min.
Maternal pain relief will be assessed with the help of visual analogue scale in the immediate postnatal period.

**Control group**

Partographic monitoring of labor was done.

Inj. tramadol 50 mg IM was used for pain relief.

Diazepam, pentazocine and drotaverine were not used.

**Assessment**

1. Duration of labor.
2. Pain relief during labor.
3. APGAR score at 1 minute and at 5 minutes.
4. Perinatal morbidity and mortality.
5. Side effect to the mother and child.

**RESULTS**

1) The rate of cervical dilatation was 1.72 cm/hour in cases and 1.23 cm/hour in controls (P <0.0001) which is significant.
2) The duration of active phase of labor was significantly reduced (3.45 hours in cases & 4.78 hours in controls, which is statistically significant P <0.0001).
3) The duration of II stage of labor was 27.4 min in cases & 34.0 min in controls (P <0.0004).
4) Majority of the patients in both groups delivered vaginally (90% in study group & 92% in controls).
5) 86% patients in study group had pain relief as compared to 22% in control group. It is statistically significant.
6) The cases had significantly more drug related side effects like nausea, vomiting, drowsiness, that subsided after 12 hours.
7) There were no significant fetal/neonatal complications in either group.

Table 1: Age distribution of subject in both groups.

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>Cases (n=50)</th>
<th>Control (n=50)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;20</td>
<td>13</td>
<td>9</td>
</tr>
<tr>
<td>21-25</td>
<td>31</td>
<td>34</td>
</tr>
<tr>
<td>26-30</td>
<td>6</td>
<td>7</td>
</tr>
<tr>
<td>&gt;30</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Maximum subjects in both groups were <25 years

Cervical dilatation was significantly faster in cases (P <0.0001)

Table 3: Duration of stages of labour.

<table>
<thead>
<tr>
<th></th>
<th>Active phase</th>
<th>II stage</th>
<th>III stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case</td>
<td>3.45 hours</td>
<td>27.4 min</td>
<td>4.46 min</td>
</tr>
<tr>
<td>Control</td>
<td>4.78 hours</td>
<td>34.0 min</td>
<td>4.45 min</td>
</tr>
</tbody>
</table>

There was statistically significant difference in the duration of II stage of labor

Table 4: Pain relief score.

<table>
<thead>
<tr>
<th>Score</th>
<th>Cases</th>
<th>Percentage</th>
<th>Control</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>1</td>
<td>02%</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>19</td>
<td>38%</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>23</td>
<td>46%</td>
<td>11</td>
<td>22%</td>
</tr>
<tr>
<td>0</td>
<td>7</td>
<td>14%</td>
<td>39</td>
<td>78%</td>
</tr>
</tbody>
</table>

Pain relief was much better in study group as compared to controls

Table 5: Complication in both group.

<table>
<thead>
<tr>
<th>Complication</th>
<th>Cases</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Vomiting</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Drowsiness</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Tachycardia</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Hypertonic contractions</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Cervical/vaginal tears</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Rupture uterus</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Side effects observed more in cases than in controls which are related to drugs

Table 6: APGAR score.

<table>
<thead>
<tr>
<th>APGAR score</th>
<th>Cases</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>At 1 min (4-5)</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>At 1 min (6-7)</td>
<td>46</td>
<td>45</td>
</tr>
<tr>
<td>At 5 min (6-7)</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>At 5 min (8-9)</td>
<td>47</td>
<td>45</td>
</tr>
</tbody>
</table>

APGAR score was comparable in both groups

**DISCUSSION**

Programmed labor protocol was designed for shorter, safer and relatively painless vaginal delivery, making it a joyful and satisfactory event.
Cervical dilatation

The cervix dilated at a faster rate (1.72 cm/hour) in programmed labor than in controls (1.23 cm/hour). It resulted in shorter labours.

According to Mishra et al.⁴ and Singh et al.,⁵ it is the effect of drotaverine. Veronica et al.⁶ noticed that rate of cervical dilatation was nearly double (2.3 cm/hour) in subjects and (1.2 cm/hour) in controls.

Duration of labour

In our study, mean duration of active phase of labor in primi was 3.45 hours (cases) and 4.78 hours (controls). It was statistically significant (P <0.0001).

Mean duration of II stage of labor was 27.4 minutes (cases) and 34 minutes (controls) which was statistically significant (P <0.0004).

Dr. Chauhan et al.⁷ found duration of first stage of labor to be 3.4 hours. Dr. Daftary et al.⁸ reported active phase duration to be 3.5 hours in cases and that of II stage of labor to be 26 minutes.

Pain relief

Pain relief plays a vital role in maternal well-being. Pain and fear retard the progress of labor. It prevents maternal hyperventilation, undue muscular efforts and exhaustion. Hence, pain relief was one of the important objectives of the study.

We observed that 43 (86%) cases had pain relief. Out of them 20 (40%) cases experienced excellent pain relief while in controls, only 22% had pain relief.

Meena Jyoti et al.⁹ noticed that 54% achieved good and 32% achieved moderate pain relief. Veronica et al.⁶ reported total pain relief in 70% cases.

Maternal outcome

None of the patients had any major complications of labour.

In programmed labor group, drug related side effects like nausea, vomiting, drowsiness, tachycardia were seen. All the side effects subsided by 12 hours after delivery.

One cases developed hypertonic uterine contractions.

Veronica et al.⁵ had similar findings. Tachycardia (80%) was the commonest side effect followed by nausea and vomiting (10%).

Neonatal outcome

APGAR score of babies in both groups were good (>7 in 94% cases & in 90% controls).

CONCLUSION

“Programmed labour protocol” provide effective analgesia and side effects of drugs which were observed are minimal, and safe for the fetus. Labour is cherished with pleasure and childbirth becomes a joyous event for the mother.

Funding: No funding sources
Conflict of interest: None declared
Ethical approval: The study was approved by the institutional ethics committee

REFERENCES


DOI: 10.5455/2320-1770.ijrcog20150434