Use of misoprostol for termination of second and third trimester pregnancy with intrauterine foetal death

Sonal Palod¹*, Trupti Nayak²

¹Department of Obstetrics & Gynaecology, PDU Medical College, Rajkot, Gujarat, India
²Department of Obstetrics & Gynaecology, MP Shah Medical College, Jamnagar, Gujarat, India

ABSTRACT

Background: Prostaglandins (PGs) is an alternative method for induction of labour in women with intrauterine foetal death (IUFD). The vaginal route is advantageous because slow peak levels, sustained for long time and fewer side effects. So this study was aimed to investigate the effectiveness of misoprostol in the induction of labour and abortion in 2nd and 3rd trimester pregnancies associated with IUFD. The objective of the study was to assess the efficacy of misoprostol in fixed dosage schedule for the termination of pregnancy in IUFD and to evaluate induction delivery interval with misoprostol.

Methods: A prospective study was conducted in department of Obstetrics & Gynaecology in tertiary care centre. Inclusion criteria included women of II or III trimester of pregnancy with IUFD. Singleton pregnancy with longitudinal lie in non-scarred uterus with confirmed intrauterine fetal death with bishop score <6. Exclusion criteria included multiple pregnancy, lie other than longitudinal, scarred uterus, hypersensitivity to misoprostol, Bishop score more than 6. Permission from IEC and written informed consent was taken before study participation. Complete examination and investigation of patient was done and induction with misoprostol was done as per dosage schedule mentioned by WHO Bellago, Italy (Feb 2007). Patient was intensively monitored intraoperatively and postoperatively. Follow-up was done till 15 days postpartum.

Results: Out of 107 cases, majority cases were of 22-28 week gestation. Maximum cases (57%) were of age group 21-25 years. Induction succeeded in 92 cases with mean induction delivery interval of 13.2 hours. 50% cases had favorable cervix with mean induction delivery interval 9.4 hours and rest had 16.53 hours ID interval with mean ID interval 13.67hours. Complete exclusion was seen in 92 cases (85.98%) and incomplete in 10 cases which required check curettage. Failure of method was seen in 5 cases (4.67%) which were terminated by alternative methods. Most common side effect was abdominal cramp (9.3%) and nausea (5.6%).

Conclusions: Vaginal misoprostol is safe and effective in termination of second and third trimester pregnancy in case of intrauterine foetal death.

Keywords: IUFD, Misoprostol, Cervical ripening, Induction delivery interval.

INTRODUCTION

A woman may need to give birth prior to the spontaneous onset of labour in situations where the fetus has died in utero (also called a stillbirth), or for the termination of pregnancy where the fetus, if born alive, would not survive or would have significant disability. This situation is psychologically stressful for the woman, her partner and family and for the health professionals caring for her.

The management of IUFD poses a dilemma. Although a significant number of these patients will spontaneously go into labour within several weeks, many do not.
Moreover, after the diagnosis, the social pressures and emotional aspects of delivery are usually considerable, and the medical consequences of postponing delivery can be significant. Within the past two decades, Prostaglandins (PGs) have provided an alternative method for induction of labour in women with IUFD. A large body of evidence exists that shows that the use of Misoprostol for labour induction is highly efficacious and safe. Misoprostol is absorbed rapidly when administered orally, vaginally, rectally or intracervically. The vaginal route is advantageous because peak levels are reached slowly and sustained for a long time and this is associated with fewer side effects. The vaginal route is also more effective than the oral route. The greater bioavailability of vaginal Misoprostol probably explains the clinical results.

The advantage of using misoprostol for induction of labour and abortion over conventional oxytocic is overwhelming. Firstly it is a stable analogue of prostaglandin E2 and thus less subjected to strict storage rules – an obvious advantage in tropical climate. Secondly, it is cheap and widely available and thirdly, in addition to intravaginal administration, it can be taken orally. These advantages make Misoprostol a favourable agent in our environment where conventional prostaglandin E2 is not only scarce but prohibitively expensive. So the aim of this study is to investigate the effectiveness of misoprostol in the induction of labour and abortion in 2nd and 3rd trimester pregnancies associated with intrauterine foetal death.

METHODS

This was a prospective study carried out from April 2011 to August 2012 at Department of Obstetrics & Gynaecology, M. P. Shah Government Medical College, Jamnagar, Gujarat.

Inclusion criteria

Women of II or III trimester of pregnancy with IUFD. Singleton pregnancy with longitudinal lie in non-scarred uterus with confirmed intrauterine fetal death with bishop score <6.

Exclusion criteria

Multiple pregnancy, lie other than longitudinal, scarred uterus, hypersensitivity to misoprostol, bishop score more than 6. All women met with inclusion criteria, gave informed written consent for participated in the study.

After admission

Complete history of the patient including her name, age, caste, residence, socio-economic status, educational status, obstetric & menstrual history were noted and history of previous uterine surgery was especially asked for. Any medical or surgical illness or drug hypersensitivity were ruled out. Specific past, personal & family history was enquired about. Thorough physical general & systemic examination was carried out. Gestational age was determined from date of last menstrual period and abdominal examination findings followed by ultrasonography. After pre-abdomen examination for determining gestational age and looking for any abdominal scar etc. local examination was done to rule out any local genital infection & cervical anomaly. This was followed by per vaginal examination for cervical length, condition of external OS.

Investigations like Hb, Blood grouping with Rh typing, urine for albumin & sugar, BT/CT, RPR, Antibody test for HIV(after pretest counselling) were done in all cases. In select-requisite cases, RBS, LFT, RFT coagulation profile are also requested. Ultrasonography was done in all cases to determine gestational age, placental localization, fetal cardiac motion localization, AFI and any congenital anomaly of fetus.

Methods for termination

After this, written informed consent of the women was taken (consent of guardian taken if the women are minor or mentally ill) explaining her condition (IUFD), methods available for termination, details of the procedure to be performed possible complications & their treatment, chances of failure and ways out.

Dose schedule to participants

Women received Misoprostol in multiple doses as per guidelines of WHO clinical guidelines Bellagio, Italy (Feb 2007).

- 13 - 17 wks: 200mcg Misoprostol PV x 6 hrly max 4 doses.
- 18 - 26 wks: 100mcg Misoprostol PV x 4-6 hrly max 4 doses.
- >26 wks: 25 - 50 mcg misoprostol 4 hrly, max 6 doses.

Failure of procedure is defined as failed expulsion of fetus at 24 hours or the occurrence of systemic adverse signs and symptoms severe enough to prohibit further use of the drug.

Examination

Vital signs i.e. T, P, BP, RR were monitored every 4 hours. Occurrence of fever, chest pain, breathing difficulty, vomiting, diarrhoea and signs of water intoxication was recorded and treated accordingly. Once having labor established, she was offered intrapartum care as per protocol for “Active management of First stage of labour” and “Active management of Second stage of labour” as and when required. PV examination was repeated every 6 hours for re-assessment. After
expulsion of fetus and placenta, fetus was examined for weight, sex & anomaly and placenta for weight, completeness and length of the cord. If required, P/S to rule out cervical injury was done followed by P/V to rule out incomplete abortion. Post expulsion sonography was done as and when required.

Completeness of abortion is defined as expulsion of both placenta and fetus without further need of Suction and Evacuation. The induction/abortion – delivery interval is defined as time of administration of 1st dose of Misoprostol to delivery/abortion.

After delivery

Antibiotic in form of Cap. Amoxicillin (500mg) 6 hourly for total of 5 days was given to all patients Inj. Anti-D 150 mcg IM was given to Rh negative mother. One tablet of cabergolin 50 mcg was given stat in case of pregnancy less than 28 wks, and two tablets stat were given in pregnancy more than 28 wks for inhibiting breast milk synthesis. Postpartum counselling was done in all women especially regarding contraception, including bereavement counselling.

Follow up

Requisite follow up was maintained and patient was called on 3rd, 7th day and then 15th day. Any significant complains like fever, abdominal pain or bleeding P/V was asked for. P/S, P/V examination was done. Post abortal advice including contraception and nutrition was provided.

RESULTS

There were 107 patients participated in study with means age of 24.61 yr ±4.38 year. 52 patients (49%) were of urban population and 51pts were of rural population. 56 pts were illiterate and 51pts were literate. The mean age of the study group was 24.61yr and the mean I.D interval was 13.20hr.

Table 1: Age wise distribution of patients and variation in induction delivery interval (ID Interval).

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>No. of cases</th>
<th>Percentage</th>
<th>Mean I.D interval (hrs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 20</td>
<td>17</td>
<td>15.88</td>
<td>13.38</td>
</tr>
<tr>
<td>21-25</td>
<td>61</td>
<td>57.01</td>
<td>13.59</td>
</tr>
<tr>
<td>26-30</td>
<td>20</td>
<td>18.71</td>
<td>11.82</td>
</tr>
<tr>
<td>&gt; 31</td>
<td>9</td>
<td>08.40</td>
<td>13.33</td>
</tr>
</tbody>
</table>

The mean age of the study group was 24.61yr and the mean I.D interval was 13.20hr.

Table 2: Parity wise distribution of patients and their variation in induction delivery interval.

<table>
<thead>
<tr>
<th>Parity</th>
<th>No. of cases</th>
<th>Mean I.D. interval (hrs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primi</td>
<td>46</td>
<td>13.5</td>
</tr>
<tr>
<td>Second gravida</td>
<td>29</td>
<td>12.43</td>
</tr>
<tr>
<td>Third gravida</td>
<td>20</td>
<td>12.3</td>
</tr>
<tr>
<td>Multigravida</td>
<td>12</td>
<td>15.45</td>
</tr>
</tbody>
</table>

Out of 107 cases, 50pts had favourable cervix and mean I.D. interval in this population was 9.41 hr. In rest 57 cases cervix was not favourable and mean I.D. interval in this population was 16.53hrs.

In our study group 18 patients belonged to 13-17wk gestation, the mean I.D. interval of this group was 14.25hr. There were 23pts in 18-21wk gestation group, the mean I.D. interval of this group was 16.34hrs. There were 34pts in 22-28wk gestation group, the mean I.D. interval of this group was 13.67hr. The rest 32pts belonged to those with gestational age > 28wks. The mean I.D. interval in this group was 9.859 hr.

In our study group, there were 36pts with uterine size between 14-20wk size and their mean I.D. interval was 15.23hr. There were 45pts with uterine size between 22-28wk and their mean I.D. interval was 13.48hrs. There were 26pts with >28wk uterine size and their mean I.D. interval was 9.23hr.

Table 3: Mean induction delivery interval in cases according to cause of intrauterine fetal death.

<table>
<thead>
<tr>
<th>Causes of IUFD</th>
<th>No. of cases</th>
<th>Mean I.D interval (hrs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abruptio placenta</td>
<td>07</td>
<td>10.64</td>
</tr>
<tr>
<td>PET/eclampsia/Chr. HT</td>
<td>18</td>
<td>12.75</td>
</tr>
<tr>
<td>Anaemia</td>
<td>03</td>
<td>15.33</td>
</tr>
<tr>
<td>Anomaly</td>
<td>12</td>
<td>20.62</td>
</tr>
<tr>
<td>Sepsis</td>
<td>19</td>
<td>13.05</td>
</tr>
<tr>
<td>Oligo/meconium/post-term pregnancy</td>
<td>05</td>
<td>10.8</td>
</tr>
<tr>
<td>Rh incompatibility</td>
<td>04</td>
<td>22.16</td>
</tr>
<tr>
<td>PROM</td>
<td>05</td>
<td>09.2</td>
</tr>
<tr>
<td>Cord pathology</td>
<td>03</td>
<td>12.5</td>
</tr>
<tr>
<td>Trauma</td>
<td>02</td>
<td>12</td>
</tr>
<tr>
<td>Idiopathic</td>
<td>29</td>
<td>10.98</td>
</tr>
</tbody>
</table>

In our study group, out of 107 cases, 92 patients had a complete expulsion without any evidence of retained products and within 24hrs. This means that 85-98% cases had successful outcome. 10 cases i.e., 9.34% cases had expulsion of fetus within 24hrs but had retained placenta for which check curettage was done. In remaining 5 (4.67%) case I.D. interval extended to be more than 24hr.
and additional help with Foley’s catheter or betadine saline induction or hysterectomy were done.

### Table 4: Outcome of cases.

<table>
<thead>
<tr>
<th>Outcome</th>
<th>No of cases</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complete expulsion</td>
<td>92</td>
<td>85.98</td>
</tr>
<tr>
<td>Incomplete expulsion</td>
<td>10</td>
<td>9.34</td>
</tr>
<tr>
<td>Failure (&gt;24 hrs)</td>
<td>5</td>
<td>4.67</td>
</tr>
</tbody>
</table>

### Table 5: Cumulative I.D. interval in the study population

<table>
<thead>
<tr>
<th>I.D interval</th>
<th>No of cases</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Within 12 hrs</td>
<td>61</td>
<td>57.0</td>
</tr>
<tr>
<td>12-24 hrs</td>
<td>41</td>
<td>38.31</td>
</tr>
<tr>
<td>&gt;24 hrs</td>
<td>5</td>
<td>4.67</td>
</tr>
</tbody>
</table>

In our study population, 61 cases i.e., 57.0% cases delivered within 12 hrs. 38.31% cases i.e. 41 patients delivered within 12-24 hrs. Only 4.67% (5 cases) cases couldn’t deliver in 24 hrs and had to be given added augmentation.

Most common side effect due to misoprostol was cramping abdominal pain seen in 10 cases i.e., 9.3%. 6 pts i.e., 5.6% had nausea and 5 patients had fever. Diarrhoea, chills and rashes were noted in 2 pts each. There was no any incidence of hemorrhage, injury, or any maternal death.

Out of 107 cases, induction of labour succeeded in 92 cases with mean induction delivery of 13.20 hours. The mean age of our study group was 24.61 years with 49% were from urban area and 54% were from rural area. 56 patients were illiterate and 51 patients were literate. 63 patients had taken antenatal care and 44 had not taken antenatal care. 76 were from lower socioeconomic class. Maximum cases were of 21-25 year age group i.e. 57% with mean induction delivery interval 13.59 hours. There were 50 cases with favourable cervix with mean induction delivery interval of 9.41hrs and 57 cases were with unfavourable cervix with mean induction delivery interval of 16.53hours.

According to weeks of gestation maximum cases were of 22-28 week gestation with mean induction delivery interval 13.67 hours. According to uterine size on admission, maximum cases were of 22-28 weeks uterus i.e. 45 cases with mean induction delivery interval 13.48 hours. Most common cause of intrauterine fetal death in our study was idiopathic followed by sepsis followed by PIH. Complete expulsion was seen in 92 cases (85.98%) and incomplete in 10 cases which required check curettage. Failure of method was seen in 5 cases i.e. 4.67%. Most common side effect seen in our study was cramping abdominal pain (9.3%) followed by Nausea in (5.6%).

### DISCUSSION

The occurrence of IUFD constitutes a major nightmare to women and attending clinicians. It is even more agonizing with a feeling of defeat to clinician if it occurs unexpectedly and the cause cannot be explained. Therefore, the ideal drug for termination of pregnancy in cases of IUFD should not only be effective and safe but should be affordable to avoid additional financial burden arising from a wasted pregnancy.

Our experience shows that misoprostol is a very effective and safe method of induction in IUFD with almost 100% vaginal delivery rate and few complications. Its stability at room temperature need for no special storage requirement and cost effectiveness make it an ideal method of induction in both developing and developed countries. The result of this study is similar to the findings in other reports.

In our study IUFD is most common in 21 to 30 year age group i.e = 57% which is similar to the study by M.O. Elmahaish et al and Jahanfar SH et al. In our study nearly 42% patients were primipara which is almost similar to the study by Jahanfar SH et al in which the incidence of IUFD in primipara was 39.2%. Among all the studies conducted, it was noted that those with favourable cervix always had lesser induction delivery interval. In our study nearly 46.7% patients had favourable cervix and hence had lesser mean induction delivery interval i.e. 9.41 hours. While 55.3% patients had unfavourable cervix and their induction delivery interval was more. This is clearly explained by the fact that those with favourable cervix were already in labour and hence actually needed only augmentation with misoprostol to clear the latent phase of labour. In our study the mean induction delivery interval is gradually decreasing with increase in weeks of gestation. This is because of increase in the receptor density in uterus as well as increase in sensitivity of uterus to the oxytocin as the pregnancy progresses (williams 23rd edition).

In our study the mean induction delivery interval (ID interval) of 14-20 wk uterine size is 15.23 hours which is almost similar to that in Abdul et al study. In our study group the most common cause of IUFD was idiopathic which is similar to Abdul et al study. The second most common cause of IUFD in our study was infection which was similar to Ezechi et al study.

In our study group 85% cases had complete expulsion and about 10% had retained products while in study of Gomez et al 25% cases had retained products.

In our study 85.9% females delivered within 24 hours which is similar to Gomez et al study. In our study nearly 5% cases delivered in >24 hrs and needed extra augmentation while in Gomez et al study 15% cases required >24 hrs to deliver and/or extra augmentation.
The most common side effects seen in our study were fever followed by diarrhoea which is similar to El Garib et al study.15

CONCLUSIONS

Results of our study suggest that vaginal misoprostol is safe and effective in termination of second and third trimester pregnancy in case of intra uterine fetal death. It is associated with very low frequency of side effects and also cost effective which indirectly increases patient compliance.

Funding: Not required
Conflict of interest: None declared
Ethical approval: The study was approved by the Institutional Ethics Committee

REFERENCES
