Introduction
Postpartum hemorrhage is a major cause of maternal death across the world (1). The diagnosis of the dramatic event of postpartum hemorrhage (PPH) requires keen clinical acumen and swift treatment. The administration of prophylactic uterotonic immediately prior to or after the placental delivery or within 1 minute of birth can reduce blood loss, thereby decreasing maternal morbidity and mortality (2).

In this regard, various uterotonic agents have been used, including oxytocin, methyl ergometrine, misoprostol, and carbetocin. Oxytocin is the most effective uterotonnic agent. The action of oxytocin depends on the number of myometrial receptors. The receptors increase as the pregnancy advances and are maximum at labor while they decrease sharply in advanced labor and the postpartum period. The receptors are minimal in number in the postpartum period, and hence the uterus becomes refractory to the action of oxytocin (3).

The best route and the dose of oxytocin is debatable. The action of oxytocin by the intravenous (IV) route is almost immediate, however, the onset of action by the intramuscular (IM) route occurs in 3-7 minutes, and the preferred route of the prophylactic of oxytocin after vaginal delivery has traditionally been IM or IV infusions with a dose of 10 IU of IM oxytocin following vaginal delivery. The difference in pre- and post-delivery hemoglobin (Hb) levels, tone of the uterus, hemodynamic changes, adverse effects of the drug, and the need for additional uterotonicics and blood transfusions were assessed based on the aim of the study.

Abstract
Objectives: Intravenous (IV) oxytocin during vaginal delivery has been rarely used since an intramuscular (IM) route or IV infusion have been preferred in this regard. The trial aimed to compare the low-dose IV bolus 3 IU of oxytocin, along with 7 IU oxytocin infusion with 10 IU oxytocin infusion in cesarean section.
Materials and Methods: A parallel control randomized study was conducted on a total of 320 consenting term pregnant women based on the inclusion criteria. The participants were randomized into either 3 IU IV bolus and 7 IU infusion of oxytocin or 10 IU IM oxytocin following vaginal delivery. The difference in pre- and post-delivery hemoglobin (Hb) levels, tone of the uterus, hemodynamic changes, adverse effects of the drug, and the need for additional uterotonicics and blood transfusions were assessed based on the aim of the study.
Results: Based on the results, more women with severe blood loss were found in the IM oxytocin group in comparison to the IV bolus with infusion group following vaginal delivery. In addition, more women had a drop in the Hb of 3 gm/dL in the IM oxytocin group compared to the IV bolus-infusion group (11% vs. 4%, odds ratio=0.768, P=0.469) although there was no statistical significance in this respect. The tone of the uterus was firmer in the IV bolus with infusion group at 3 and 5 minutes. Eventually, the difference in hemodynamic changes, side effects, and the need for additional uterotonicics or blood transfusions was not significant.
Conclusions: In general, an IV bolus of 3 IU with a 7 IU infusion of oxytocin is as safe as and more effective than the IM injection of 10 IU of oxytocin at the time of vaginal delivery for the prevention of postpartum hemorrhage.
Keywords: Oxytocin, Intravenous bolus, Intramuscular, Blood loss, Hemodynamics, Postpartum hemorrhage

Randomised Control Trial of 3 IU Intravenous Oxytocin Bolus With 7 IU Oxytocin Infusion Versus 10 IU Intramuscular Oxytocin in the Third Stage of Labour in the Prevention of Postpartum Hemorrhage

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Intravenous oxytocin in the dose of 3 IU along with 7 IU infusion is as safe as and more effective than IM oxytocin when used post vaginal delivery.

Concerns of hemodynamic instability after the administration of intravenous bolus oxytocin were not experienced in this study.

Materials and Methods

A parallel control randomized study was performed on 320 consenting women from 25th February to 25th May, 2020.

The present study included 320 pregnant women of term gestation visiting the Department of Obstetrics and Gynecology, Shri B.M. Patil Medical College Hospital and Research Center, Vijayapura. Written informed consent was obtained from all participants. Complete history and clinical findings were recorded in the prescribed format. In addition, the complete blood count was done for all patients on admission to the labor ward. The test was repeated after 48 hours of delivery. Participants were assigned to randomized groups according to a computer-generated randomization chart obtained from the web site of www.Randomization.com.

The study comprised of two groups (groups I and II). Group I (n = 160) received 3 IU IV oxytocin bolus with 7 IU oxytocin in IV infusion. Group II (n = 160) received 10 IU IM oxytocin.

Group I: 3 IU of oxytocin was diluted and administered intravenously over 1 minute. The drug was given at the time of delivery of the anterior shoulder of the baby. It should be noted that 1 mL oxytocin equals 5 IU. This was diluted in 4 mL of 0.9% normal saline, making it 1 IU per mL, and 3 mL was given bolus over 1 minute. The remaining 2 mL was given in IV infusion, along with another 5 IU of oxytocin making it 7 IU. The IV fluid was either Ringer’s Lactate or 0.9% normal saline.

Group II: 10 IU IM oxytocin was given to the mother during the delivery of the anterior shoulder of the baby. Two ampoules of 5 IU of oxytocin were loaded in a single syringe and administered in the deltoid region.

The participants who were augmented with oxytocin during the first stage of labor were not excluded as the active management of the first stage of labor was followed in the institution.

The primary outcome was to assess the blood loss by calculating the difference in the pre- and post-delivery hemoglobin (Hb) levels. The Hb levels were estimated as part of the complete blood count. The first pre-delivery complete blood count and post-delivery complete blood count were conducted 48 hours after the delivery.

The secondary measured outcomes were changes in the heart rate and systolic and diastolic blood pressure (BP), the tone of the uterus, the need for extra uterotonics and blood transfusion, and the side effects of the drugs, including nausea, vomiting, and the like.

The tone of the uterus was assessed on a 5-point scale. The uterine tone was assigned as 1 if atonic, 2 if there were partial, inadequate uterine contractions, 3 if there were adequate contractions, 4 if the uterus was well contracted, and 5 if the uterus was very well contracted.

Sample Size Calculation

Based on the study by Charles, Anger, Dabash et al (9), 320 (160 per group) pregnant women delivering vaginally are required to have a 90% chance of detecting, as significant at the 5% level, a decrease in the mean blood loss (mL) from 180 ± 100 in group I to 214±100 in group II. The calculation was based on the following formula:

\[ n = f(\alpha/2, \beta) \times 2 \times \sigma^2 / (\mu_1 - \mu_2)^2 \]

where \( \mu_1 \) and \( \mu_2 \) are the mean outcomes in the study groups, respectively, and \( \sigma \) denotes the standard deviation.

Data Analysis

The statistical analysis of all characteristics is summarized descriptively. The summary statistics of N, mean, standard deviation (SD) will be used for continuous variables. Regarding categorical data, the number and percentage are applied in data summaries. Further, data are analyzed by the Chi-square test for association, comparison of means using a t test, ANOVA, and diagrammatic presentation. T-test/Z-test is used to know the efficacy of each method and the chi-square test is employed to compare the efficacy between the groups.

Results

During the study period (from 25th February to 25th May...
2020), a total of 1,481 women delivered, of which 920 cases underwent C-section and 561 delivered vaginally. Of women who delivered vaginally, 506, 21, and 34 cases were term deliveries, vaginal birth after caesarean, and preterm deliveries, respectively. Of 506 women who delivered at term, 350 women met the inclusion criteria. Out of these 350 pregnant women, 320 gave their consent by signing an informed and written consent form and were included in the study.

Based on the results, there was no statistical difference regarding the age of the participants of groups I and II (23.61 vs. 23.87, \( P = 0.482 \)) (Table 1). The weight of the participants in both groups was matched (60.31 vs. 61.31, \( P = 0.74 \)), and the number of primigravidae and multigravida were equally matched in both groups (Table 2). The difference in pre- and post-delivery Hb values was calculated as well. In 90.6% and 88.1% of the women of groups I and II, the difference in the pre- and post-delivery Hb was less than 3 g/dL. A difference in the Hb values of more than 3 g/dL was observed in 9.4% and 11.9% of the participants of groups I and II, respectively. There was an increased risk of excessive blood loss in group II in comparison to group I (OR 0.768, 95% CI: 0.375-1.57) (Table 3). Although the difference was not statistically significant, women in group II lost more blood. The number of the given episiotomies, incurred cervical lacerations, and observed perineal tears were similar in both groups. Three patients in group II and three patients in group I required additional uterotonics and styptics, respectively. There was only one blood transfusion in group II.

The majority of participants of group I reached grade 5 of the tone of the uterus at 3, 5, and 10 \( (P \text{ value at } 3 \text{ minutes } = 0.025, P \text{ value at 5 minutes } = 0.043) \) minutes, implying that the IV bolus oxytocin, along with the infusion acted faster on the uterus (Figure 1).

No statistically significant differences were found in the heart rate, systolic BP, and diastolic BP before the intervention and at 5, 10, and 60 minutes in both groups (Figure 2).

**Discussion**

The study aimed at comparing the efficacy of bolus oxytocin with IM oxytocin regarding preventing excessive bleeding in women delivering vaginally. The study included 320 women who were randomized into two groups each including 160 women. Women in groups I and II were matched equally in terms of age, parity, and weight. Our results demonstrated that there was no difference in the two groups as the Hb drop was less than 3 gm/dL. However, the chance of bleeding was more in group II as compared to group I if the drop in Hb was more than 3 g/dL (OR = 0.768, 95% CI: 0.375-1.57). In a study conducted in the Republic of Ireland (8),

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Group I</th>
<th>Group II</th>
<th>( P ) Value</th>
<th>95% CI of Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>23.61±3.33</td>
<td>23.87±3.35</td>
<td>0.482</td>
<td>-1.00</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>60.31±8.19</td>
<td>61.31±9.41</td>
<td>0.313</td>
<td>-2.94</td>
</tr>
</tbody>
</table>

Note. SD: Standard deviation; CI: Confidence interval; SE: standard of error.

The table below shows the comparison of parity, Hb percentage, traumatic features, additional applied drugs, and the required blood transfusion between the two groups.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Group I</th>
<th>Group II</th>
<th>Chi-square</th>
<th>( P ) Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gravida</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primigravida</td>
<td>62</td>
<td>54</td>
<td>0.865</td>
<td>0.352</td>
</tr>
<tr>
<td>Multigravida</td>
<td>98</td>
<td>106</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes in pre- and post-delivery Hb (g/dL)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;3</td>
<td>145</td>
<td>141</td>
<td>0.527</td>
<td>0.468</td>
</tr>
<tr>
<td>&gt;3</td>
<td>15</td>
<td>19</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Episiotomy</td>
<td>99</td>
<td>95</td>
<td>0.209</td>
<td>0.647</td>
</tr>
<tr>
<td>Cervical lacerations</td>
<td>3</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perineal tears</td>
<td>6</td>
<td>2</td>
<td>2.051</td>
<td>0.152</td>
</tr>
<tr>
<td>Tachysystole</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Additional used uterotonics</td>
<td>0</td>
<td>3</td>
<td>4.013</td>
<td>0.134</td>
</tr>
<tr>
<td>Applied styptics</td>
<td>3</td>
<td>1</td>
<td>1.01</td>
<td>0.314</td>
</tr>
<tr>
<td>Any blood transfusion</td>
<td>0</td>
<td>1</td>
<td>2.013</td>
<td>0.366</td>
</tr>
<tr>
<td>Total</td>
<td>160</td>
<td>160</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note. Hb: hemoglobin.

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**Table 1. Association of Age and Weight With Groups I and II**

**Table 2. Comparison of Parity, Hb Percentage, Traumatic Features, Additional Applied Drugs, and the Required Blood Transfusion Between the Two Groups**
10 IU IV oxytocin was administered over 1 minute and compared to 10 IU of the IM administration of oxytocin after vaginal delivery. Based on the results, although no difference in the prevention of PPH was observed in mild PPH, it was evident in severe PPH, and admission to high dependency units was lesser in the IV bolus group in comparison with the IM group (9). In another large-scale study conducted in Egypt, 10 IU IV bolus oxytocin was found to be superior to 10 IU IM oxytocin in vaginal delivery (9). However, the other systematic review could not establish whether the IV or IM route was superior and concluded that larger randomized control trials were required in this regard (10).

The hemodynamics of the patients were stable in both groups and no difference was found between the groups in this regard. The results represented that IV bolus oxytocin in the dose of 3 IU was as safe as IM oxytocin, which is in line with the findings of other recent studies on IV bolus oxytocin in post-delivery cases (8,9). It also shows that the older concept that IV oxytocin is unsafe and may cause hemodynamic instability does not hold well in cases of vaginal delivery (9). Similar studies conducted administering IV bolus oxytocin showed no side effects or adverse outcomes in comparison to IV oxytocin infusion (11,12). Some studies reported that there were stronger uterine contractions followed by IV bolus oxytocin, but this route was not frequently used due to an immediate decrease in the BP, even though this side effect of IV bolus oxytocin was frequently observed in patients who underwent caesarean section under general anesthesia (5,13). Various studies showed that IV bolus oxytocin, when given in large doses, has adverse effects such as a decrease in BP, headache, nausea, vomiting, and even myocardial infarction in extreme cases. However, in our study, a smaller dose of IV oxytocin when given in bolus demonstrated no serious hemodynamic changes as seen in other studies (5,14,15). According to a Syntocinon Infusion trial, hemodynamic changes occurred mainly due to the regional anesthesia during caesarean section instead of IV bolus oxytocin (16).

In a study conducted on women undergoing C-section, the IV bolus of either 2 IU or 5 IU, along with the infusion of 0.25 IU/mL was compared to the IV infusion of oxytocin of 0.5 IU/mL. Based on the results, there were marked hemodynamic changes in heart rate and mean arterial pressure in the bolus-infusion groups compared to the only infusion group. However, changes were less in the 2 IU bolus group. The uterine tone increased in the bolus-infusion group in comparison to the infusion group (4). In the present study, the uterine tone was assessed to be firmer in group I than in group II at 3 and 5 minutes. This would hypothetically reduce the need for additional uterotonics. However, this did not reflect in the results thus further large-scale research would be needed in this respect. Another study attempted to calculate the least bolus dose of oxytocin that could be required in scheduled C-sections and for C-sections done for difficult labor. It was recommended that the 1 IU of oxytocin for elective C-section and 1-1.5 IU bolus oxytocin for cesareans done for labor progress should be administered in this regard (17). The World Health Organization advises 10 IU oxytocin either IV or IM for the prevention of postpartum hemorrhage (18). Most of the recommendations for postpartum hemorrhage do

Note. OR: Odds ratio; Hb: Hemoglobin; CI: Confidence interval.

Table 3. OR of Pre- and Post-delivery Hb g/dL in Group II Compared to Group I

<table>
<thead>
<tr>
<th>Hb Changes (g/dL)</th>
<th>OR</th>
<th>P Value</th>
<th>95% CI OR</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;3</td>
<td>1.303</td>
<td>0.469</td>
<td>0.637</td>
</tr>
<tr>
<td>&gt;3</td>
<td>0.768</td>
<td>0.469</td>
<td>0.375</td>
</tr>
</tbody>
</table>

Figure 1. Comparison of Tone of Uterus at 1, 3, 5, and 10 Minute(s) Between the Two Groups.

Figure 2. Comparison of Hemodynamic Parameters Between the Two Groups. Note: SBP: Systolic blood pressure; DBP: Diastolic blood pressure.
not differentiate between vaginal delivery and C-section. Similarly, there are no guidelines to suggest different doses for women undergoing an elective C-section and women who are in labor. IV bolus oxytocin has been studied in various doses of 2, 3, 5, and 10 IU during C-section (4-7). Studies on the bolus-infusion of oxytocin during vaginal delivery are necessary. This route could be a preferred route in cases where IM injections are contraindicated like thrombocytopenia.

Limitations of the Study
The study did not record the visible blood loss at labor and did not include the fetal outcome as the drug was administered after the delivery of the fetus.

Conclusions
Overall, 3 IU IV oxytocin bolus, along with 7 IU oxytocin infusion reduced the occurrence of severe postpartum hemorrhage in comparison to 10 IU IM oxytocin without causing any significant hemodynamic changes. Finally, the uterine tone increased faster in women who were administered IV bolus oxytocin and infusion as compared to IM oxytocin.

Authors’ Contribution
AMB and RGY conceptualized, designed, executed the study and prepared the manuscript. SSK, SSM and SSS executed the study. SRM oversaw the study and managed the logistics of the study.

Conflict of Interests
Authors declare that they have no conflict of interests.

Ethical Issues
The study was approved by the Institutional Ethical Committee of BLDE University (No. BLDE(DU)/IEC/355/-2019-20) and registered in the Clinical Trials of India (Ref No. CTRI/2020/02/023521).

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References