Does lidocaine gel produce an effective analgesia prior to copper IUD insertion? Randomized clinical trial

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ABSTRACT

Background: IUD is a small contraceptive device, often containing either copper or levonorgestrel, which is inserted into the uterus. Objective of present study was to determine if lidocaine gel prior to intrauterine device (IUD) insertion decreases pain with the insertion procedure among multiparous women choosing the copper T380A-IUD.

Methods: It is a randomized double-blind controlled trial carried out at Assiut Women's Health Hospital, Assiut, Egypt. Parous women eligible for Copper IUD insertion attended the Family Planning Clinic were recruited and randomized in a 1:1 ratio to lidocaine gel or placebo. Two ml of the study medications were topically placed on the cervix 3 minutes before IUD insertion. The primary outcome was the difference in pain scores using a 10-cm Visual Analogue Scale (VAS) during IUD insertion. We considered a 1.5 cm difference in VAS scores between study groups as clinically significant.

Results: One hundred women consented to participate and randomized either to group I: lidocaine group or group II: placebo group. Both groups were homogenous in baseline socio-demographic data. There was significant difference in mean pain scores for IUD placement between women who received lidocaine gel and placebo at two steps of insertion (at vulsellum application and at uterine sounding) while the rest of steps show no statistical significant difference (p>0.000). There were no statistical significant differences between both group as regard the ease of insertion, the duration of insertion and the satisfaction score after the procedure (p>0.05).

Conclusions: This study depicts that the use of lidocaine gel prior to copper IUD insertion in multiparous women could partially reduce the pain during tenaculum placement and uterine sounding.

Keywords: Contraception, Intrauterine device, Lidocaine, Pain relief

INTRODUCTION

IUD is a small contraceptive device, often containing either copper or levonorgestrel, which is inserted into the uterus. They are one form of long-acting reversible contraception (LARC) which is the most effective types of reversible birth control. Depending on the country, the use of IUDs worldwide ranges from 2% to 75%. On average, 15% of reproductive-aged women in developing regions and 9% in developed regions use IUDs. IUDs are considered appropriate for most women, including nulliparous women and adolescents.

Many studies were published about the use of different medications or technical modifications in the insertion procedure trying to minimize the pain during IUD insertion. These include non-steroid anti-inflammatory drugs (NSAIDs), misoprostol for pre-insertion cervical ripening, nitric oxide donors, and local anesthetics.
Technical modifications were also published in order to omit or modify some steps during the insertion to decrease the participants’ pain perception.11-14

The levels of pain that women experience during IUD insertion vary in published reports. Most women experience mild to moderate discomfort during IUD insertion. Rarely, the pain is severe and associated with nausea and weakness.6 Predictors of pain during IUC insertion include nulliparity, age greater than 30 years, a longer interval since last pregnancy or menses, history of dysmenorrhea, and not currently breastfeeding.5

Lidocaine gel is routinely used in the distal urethra prior to Foley catheter placement, as well as in the nasal canal prior to nasogastric or nasotracheal tube placement. In each of these cases lidocaine gel has been shown to decrease pain scores associated with insertion.15 The endocervical canal is lined with columnar epithelium as is the nasal canal. The ecto-cervix and distal urethra stratified squamous epithelium lending biologic plausibility to this intervention for both tenaculum placement to the ectocervix as well as IUD insertion.

Therefore, the current study aims to evaluate the analgesic effect of topical lidocaine gel placed on the cervix prior to IUD insertion.

METHODS

The current study was a randomized, double-blind trial. The study was conducted Assiut Women Health Hospital, Egypt between October 2016 and March 2017. The Institutional Ethical Review Board approved the study, and we obtained a written informed consent from all participants before enrollment.

A total of 100 women attending the family planning outpatient clinic seeking IUD for contraception were invited to participate in our study. We included in our study women aged (18-50 years) not taken analgesics or anxiolytics in the 24 hours prior insertion and not taken misoprostol prior to insertion.

A statistician, not otherwise involved in the study, prepared a computer generated random table and placed the allocation data in sequentially numbered opaque sealed envelopes. Each envelope had a card noting the group identifier inside. A single pharmacist was responsible packaging of both gels in sterile tubes with labeling them as A and B. Only the Pharmacist knew what the medication in tube A and B was, so neither the clinician nor the women knew the type of the preparation.

The participated women were entered the screening phase of the study. This phase included history taking (about age, parity, mode of previous delivery, pattern of menstrual cycle, history of pelvic surgery, sexually transmitted diseases, interval from last pregnancy or last menses or last miscarriage and whether they had taken any pain or anxiety medications that day).

Before insertion, one of the study researchers explained the standard 10-cm visual analog scale (VAS) to the participants for pain scoring.16 The severity of pain was assessed with VAS (with 0= no pain and 10= worst imaginable pain). Each woman received a copper T380A IUD (Pregna®T380A; Pregna International Ltd USA, Mumbai, India) for insertion.

The eligible women were allocated to either group I (lidocaine group) 1 mL lidocaine gel was placed on the cervix at the anterior lip and 1 mL in the cervix up to the level of the internal os using cotton swab or group II (placebo group) with an inert gel similar in appearance, color and consistency. The placebo cream was manufactured in the Department of Pharmaceuticals-Faculty of Pharmacy. After a three-minute waiting period, the IUD was inserted in the standard fashion.

The IUD was inserted by one of the study investigators. Firstly, the speculum was placed into the vagina and the cervix was cleansed with Povidone iodine. Then, traction was applied on the cervix using tenaculum and the uterine sound was introduced followed by the IUD insertion. Immediate complications as uterine perforation and vasovagal attack in addition to the duration of insertion were recorded. A research assistant asked the women to rate the intensity of pain at five consecutive steps; at speculum placement, at tenaculum placement, at sound insertion, at IUD insertion and 5 minutes after the end of insertion using the same 10-point VAS.

After the end of procedure, the clinician assessed the ease of IUD insertion using the ease of insertion score (ES). The ES is a graduated VAS-like scale from zero to 10; in which 10 means terribly difficult insertion and zero means very easy insertion.

At 15 minutes post insertion; all women were asked to report their level of satisfaction with IUD insertion by completing a 10-cm VAS (with 0= no satisfaction and 10= maximum satisfaction).

The primary outcome was the difference in mean pain VAS scores during IUD insertion. The secondary outcomes included the mean pain scores during speculum, tenaculum placement, sound insertion and 5 minutes post-insertion, the ease of insertion score, the women’s satisfaction score and the duration of insertion.

Statistical analysis

The data were collected and entered into a Microsoft Access database and were analyzed using the Statistical Package for Social Science (SPSS Inc., Chicago, version 21). The demographic characteristics and baseline data were compared between the groups. The outcome variables were calculated using Student’s t test. For
dichotomous variables, chi-square was used to estimate the significance value. For analysis, p<0.05 was considered to be significant.

RESULTS

One hundred and twenty women were counselled for participation, however; 20 women were excluded (8 patients were not willing to share in the RCT and 12 women did not meet the inclusion criteria).

One hundred women consented to participate and randomized either to group I: lidocaine group or group II: placebo group. Both groups were homogenous in baseline socio-demographic data without significant differences (Table 1).

Table 1: Socio-demographic criteria of the study participants.

|                  | Lidocaine group (n=50) | Placebo group (n=50) | P-value |
|------------------|------------------------|----------------------|---------|
| Age              | Mean±SD                | 30.58±5.28           | 30.00±7.76 | 0.747 |
| Parity           | Mean±SD                | 2.72±0.86            | 2.56±1.30 | 0.371 |
| Residency        |                        |                      |          |
| Urban            | 28 (56%)               | 29 (58%)             |          |
| Rural            | 22 (44%)               | 21 (42%)             | 0.316   |
| Previous VD      |                        |                      |          |
| Yes              | 37 (74%)               | 38 (76%)             | 0.649   |
| No               | 13 (26%)               | 12 (24%)             |          |
| Previous CS      |                        |                      |          |
| Yes              | 28 (58%)               | 25 (50%)             | 0.705   |
| No               | 22 (44%)               | 25 (50%)             |          |
| Previous miscarriage |                |                      |          |
| Yes              | 18 (36%)               | 12 (24%)             | 0.423   |
| No               | 32 (64%)               | 38 (76%)             |          |

The mean satisfaction scores were 4.72±0.45 and 4.82±0.39 in the lidocaine and placebo groups respectively (p=0.979). Additionally, no difference in the ease of insertion score was observed among both groups (p=0.242). Additionally, the duration of IUD insertion was similar in both groups (p=0.125). No cases of uterine perforation or vasovagal reactions were observed in both groups (Table 3).

DISCUSSION

Pain during IUD insertion is multifactorial; application of the tenaculum on the cervical lip can induce severe pain. In addition, insertion of sound and IUD inside the uterine cavity can add to pain perception. Transmission of pain from the uterus occurs through two different visceral pain pathways: parasympathetic (S2–S4) provides sensory innervation to the cervix and lower portion of the uterus and sympathetic (T10–L1) provides sensory innervation to the fundus.

Local anesthetics used prior to IUD insertion include a number of formulations (e.g., gel, injections and spray) and different techniques for administration (intracervical and paracervical). They have several advantages; however, no studies have provided strong evidence that various lidocaine formulations provide significant pain relief for women undergoing IUD insertion.

Lidocaine has a rapid onset of action, reportedly around 2 minutes or less. As with duration of action, onset of action is liable to vary among application sites. We allowed 3 minutes to elapse between administration of lidocaine and IUD insertion, and based on the results, this time period was sufficient for the drug to take effect.

Our results showed significantly lower pain scores during tenaculum placement as compared with the placebo group (p=0.000) and sounding during IUD insertion in
the lidocaine group as compared with the placebo group (p=0.000). Oloto et al evaluated the efficacy of 2% lignocaine gel, inactive placebo gel or ‘no treatment’ applied to the cervical canal for reduction in pain in nulliparous or parous women. Intracervical application of 2% lignocaine gel resulted in a significant reduction in pain compared with no active treatment (placebo gel or no treatment; p=0.025). In contrast to Maguire et al study, Pain (measured by 100 mm VAS) was greatest during sounding and was similar between groups (51.6 mm in the placebo group versus 55.5 mm in the lignocaine group; p=0.33). Additionally, no significant difference between both groups for pain scores during IUD insertion (p=0.28).

In McNicholas et al, study, no significant difference was observed between lidocaine and placebo gels at either of the time points studied for the overall participants, nor in the individual subgroups of nulliparous and parous women. The study groups did not differ significantly in median pain scores at tenaculum placement (p=0.54 for nullipara and 0.23 for multipara) or during IUD insertion (p=0.18 for nullipara and 0.72 in multipara).

Similar to present results, Torky et al reported that topical lidocaine gel and spray significantly reduce the pain induced by vulsellum application (p=0.003) with no affection of the pain perception during IUD insertion (p=0.059). However, they didn’t assess neither the ease of insertion score rated by the insertion physician or the participants’ satisfaction score.

Fouda et al reported that the use of oral diclofenac potassium combined with 2% lidocaine gel slightly reduced pain scores during tenaculum application and IUD insertion. On the contrary, Abbas et al reported in their study on oral diclofenac potassium plus lidocaine cream significantly reduction of the VAS pain scores during injection of the dye and up to 30 minutes post-procedure with p=0.0001. No significant differences in VAS score after speculum or tenaculum placement.

Karasu et al reported that lidocaine spray is effective for reducing the pain experienced during IUD insertion. It reduces pain related to both tenaculum use (p<0.001) and IUD insertion (p<0.001). However, in this study the researchers did not standardize all other confounding factors that could affect the effect of lidocaine on pain perception therefore, their results could be biased.

The strengths of our study include that it was a double-blind randomized controlled trial with neither women nor the clinicians being aware of the group assignment. The study had its limitations including that the study focused on one type of IUD because the levonorgestrel IUD is not widely used in Egypt due to its high cost, and consequently the data are applied only for the copper IUD. A second limitation was the subjectivity in reporting pain through VAS score, as there is no objective parameters to evaluate pain. Furthermore, none of the included women were nulliparous because IUD insertion is not requested by this group in Egypt.

CONCLUSION

In conclusion, topical application of lidocaine gel on the cervix prior to copper T380A IUD insertion showed significantly lower pain scores only during tenaculum placement and sounding.

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