Treatment of irregular bleeding with oestradiol during long-term levonorgestrel-releasing intrauterine system (LNG-IUS) use

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Abstract

Objectives: Up to 60% of women remove their levonorgestrel-releasing Intrauterine System (LNG-IUS) prematurely, mostly because of irregular bleeding disturbances up to six months after insertion. Assuming ongoing bleeding is due to atrophic bleeding of the endometrium, treatment with oestrogen could be effective. No study in literature supports oestrogen therapy for irregular bleeding in LNG-IUS users.

Study design: A prospective cohort study was conducted in two hospitals to evaluate the effect of oral oestrogen (started six months after LNG-IUS insertion) on irregular bleeding days. The participating women received 2mg oestradiol daily for six weeks. We observed the discontinuation rate, the occurrence of adverse events and side effects due to oestrogen. Women completed questionnaires at baseline and three months after start of the treatment.

Results: Nineteen women were included. A median of 21 irregular bleeding days per month at baseline decreased to a median of five irregular bleeding days per month (p=0.003) after the use of oestradiol for six weeks. Three out of nineteen women had their LNG-IUS removed at three months follow-up because of persistent irregular bleeding. No adverse events were reported, however, 68% of the participants reported side effects.

Conclusion: In this study we observed a decrease in the number of irregular bleeding days in LNG-IUS users with ongoing irregular bleeding (>6 months after insertion), after administering oral oestradiol for six weeks. We propose to conduct a larger prospective trial to evaluate oestradiol as treatment option for irregular bleeding.

Introduction

An intrauterine device is one of the most common methods for contraception used by women (between 15-49 years) all over the world: in 2015 fourteen per cent used an intra uterine contraceptive device [1]. One of the most commonly used intrauterine devices is the levonorgestrel intrauterine System (LNG-IUS; Mirena®, Bayer HealthCare, Pittsburgh, PA, USA), which initially releases 20 micrograms levonorgestrel intrauterine every 24 hours. In the following five years, this progesterone declines to a rate of fourteen micrograms per day, which is proven still clinically effective for contraception. The LNG-IUS is also approved as a treatment option for heavy menstrual bleeding [2]. However, one of the main reasons to remove the LNG-IUS prematurely is irregular bleeding disturbance. The incidence of irregular bleeding decreases over the first six months of use [3]. In some cases, irregular bleeding may continue after six months or occurs de novo over time during LNG-IUS use. The guidelines of the National Institute for Health and Clinical Excellence (NICE) make healthcare professionals aware of the fact that up to 60% of women remove the LNG-IUS prematurely [3]. The Dutch general practitioners’ guideline recommend treatment with oral oestrogen in women with ongoing irregular bleeding more than six months after insertion of their LNG-IUS [4]. However, in literature, not one study supports oestrogen therapy for irregular bleeding in LNG-IUS users [5]. Because of the physiological effect of levonorgestrel in downregulation of oestrogen receptors in the endometrium resulting in endometrial atrophy, we hypothesize that the addition of oestradiol might be beneficial for regeneration of the endometrium and consequently decreasing irregular bleeding [6,7].

In this prospective cohort study, we evaluate the effect of oral oestradiol on ongoing irregular bleeding disturbances during LNG-IUS use more than six months after insertion.

Material and methods

This multicentre prospective cohort study was conducted between May 2017 and January 2018 in two teaching hospitals in the Netherlands.

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Women with an LNG-IUS in situ for more than six months, who visited the outpatient clinic with ongoing or de novo irregular bleeding disturbances were included. Exclusion criteria were women younger than 18 years, malposition of the LNG-IUS, known abnormal cervical cytology or (pre-) malignancies of the uterus, the presence of polyps, myomas on ultrasound or a contra-indication for oestradiol (history of thrombosis or mamma carcinoma). The study was registered in the Dutch trial register as NL8007. The ethics committee of the hospital approved the study on 12 May 2017 (N17.070). Written informed consent was obtained from all patients.

Patient characteristics including age, BMI, parity, indication for the LNG-IUS, bleeding pattern, the number of bleeding days, satisfaction with the bleeding pattern and previous therapies for the current complaints were collected at baseline.

Menstrual blood loss was assessed using a menstrual bleeding chart. The definitions used in this chart are described in table 1 [8]. Transvaginal ultrasound was performed to collect data about the endometrial thickness, the position of the LNG-IUS and to check for intrauterine abnormalities. All women used 2 mg oestradiol orally once a day for six weeks. Questionnaires were completed at baseline and three months after start of treatment including a menstrual bleeding chart. The questionnaires included the following themes: therapy compliance, the bleeding pattern, number of bleeding days, satisfaction with the bleeding pattern (on a five-point Likert scale), satisfaction with the treatment, side effect of the treatment and adverse events. Additionally, patients were asked whether the LNG-IUS was removed. Data were collected in a Statistical Package for the Social Sciences-database, in which individual study numbers were assigned to every patient, ensuring patient anonymity. The categorical data were expressed as numbers and percentages. Continuous data with normal distribution were expressed as mean ± standard deviation, while abnormal distributed continuous data were expressed as median and interquartile ranges. Wilcoxon signed rank test was used to compare continuous variables. A P-value <0.05 was considered statistically significant.

Results

A total of 25 women were included initially, one woman did not start with oestradiol after reading the warning on adverse effects as described in the leaflet of the oestradiol. Five women were lost to follow up despite to frequent recalls. In total nineteen women completed the follow-up period of three months and were used for analysis. Baseline characteristics are shown in table 2. The mean age was 30.2 (± 8.5) years. Most women initiated the LNG-IUS for contraception (57.9%).

Fourteen out of the 19 patients used oestradiol 2 mg for six weeks as prescribed (range 28–90 days). One patient used the oestradiol for four weeks because she was already satisfied with her menstruation pattern, and four patients used the oestradiol for a longer time up to 90 days. The total number of bleeding days decreased after treatment with oestradiol (Table 3). The baseline median of 21 (IQR 16.0-30.0) irregular bleeding days per month decreased to a median of 5 (IQR 3.0-11.0) irregular bleeding days per month (p=0.003). Before treatment with oestradiol, 68% of the woman reported spotting in their menstruation pattern, which decreased to 32% of the woman after treatment with oestradiol. Furthermore, 42% of the women reported normal menstruation after treatment with oestradiol.

Three women (15.8%) removed their LNG-IUS prematurely during the follow-up period, two women because of persisting irregular bleeding and one woman because of abdominal pain. No adverse effects were reported. Side-effects were reported by 68.4% of women. Most reported side-effects included painful swollen breasts (26%), mood swings (21%), headache (16%), weight gain (16%) and vaginal complaints (11%).

13 of the 19 (68.4%) women are more satisfied on a five-point Likert scale after the treatment with oestradiol with their bleeding pattern (Figure 1). 29% of women were (very) unsatisfied, 28% or the bleeding pattern and intervals (time between two bleeding <20 days). The definitions used in this chart are described in table 1 [8].

Table 2. Baseline characteristics. *2 missing; a: Data are given as mean ± standard deviation.

| Characteristics                  | N=19 |          |
|----------------------------------|------|----------|
| Maternal age, y*                 | 30.2 (± 8.7) |          |
| Body Mass Index*, kg/m²a         | 23.4 (± 3.0) |          |
| Parity, n (%)                    | 9 (47.4) |          |
| Nulliparity                      | 10 (52.6) |          |
| Multiparity                      | 11 (57.9) |          |
| Indication of LNG-IUS, n (%)     | Contraception   | 5 (26.3) |
|                                  | Heavy menstrual bleeding | 2 (10.5) |
| Spotting during use of OAC       | 1 (5.3) |          |
| Satisfaction bleeding pattern, n (%) | 1 very unsatisfied | 6 (31.6) |
|                                  | 2 unsatisfied | 11 (57.9) |
|                                  | 3 neutral | 1 (5.3) |
|                                  | 4 satisfied | 0 |
|                                  | 5 very satisfied | 1 (5.3) |
| Previous treatment, n (%)        | No | 13 (68.4) |
|                                  | Yes, lynestrenol | 6 (31.6) |

Table 1. Definition of the menstrual bleeding chart [8].

| Normal menstruation | Cycle of 24-38 days, menstrual period 4, 5-8 days |
| Heavy menstrual bleeding | Regular cycle, severe and/or prolonged blood loss (≥ 2 days of clots and/or menstrual duration>8 days) |
| Metrorrhagia | Unrecognizable cycle, varying amounts and intervals (time between two bleeding <20 days) |
| Intermenstrual bleeding | Normal menstruation with intermenstrual bleeding. |
| Contact bleeding | Blood loss as a result of coitus |
| Spotting with hormonal therapy | Slight blood loss between (or without) the withdrawal bleeding or natural periods. |
| Blood loss corresponding a section-niche | >2 days of brown discharge at the end of the menstruation (duration>8 days) or intermenstrual bleeding that starts within 5 days after menstruation. |

Figure 1. Satisfaction blood loss before and after treatment with oestradiol
women scored neutral and 45% of the women were (very) satisfied with the oestradiol treatment.

Discussion and conclusion

In this feasibility study, a reduction in the number of bleeding days during long term LNG-IUS use was found after treatment with oestradiol for six weeks. No adverse effects were reported, but several side-effects were reported (i.e. mood swings and swollen breasts). This suggests that oestradiol might be an effective treatment for ongoing irregular bleeding disturbances in long-term LNG-IUS users.

Hypothetically, the irregular bleeding is caused by the progesterone-only treatment in the LNG-IUS. Progesterone causes an increase of fragile superficial vessels of the endometrium. These vessels tear down easily, resulting in a breakthrough bleeding [9]. The progesterone in the uterus has a mechanism of downregulating oestrone receptors in the stroma cells of the basal layer of the endometrium, which leads to endometrial atrophy [6] By addition of oral oestrone, an up-regulation of oestrone receptors will take place, which might have a positive effect on irregular bleeding. Evidence for above hypothesis is lacking. After systematic search in literature we did not find publications on treatment for ongoing or the novo irregular bleeding disturbances more than six months after LNG-IUS insertion [10-13]. We found four studies describing prophylactic treatment regimens, starting straight treatment after LNG-IUS insertion.

One of the studies evaluated the prophylactic effect of oestradiol on irregular bleeding after insertion of the LNG-IUS [7]. However, a disadvantageous effect of oestradiol patches was shown. We hypothesize that there is a different mechanism of irregular bleeding disturbances in short-term use compared to long-term use. Probably the atrophic endometrium bleeding occurs later on and oestrogen treatment straight after insertion results in a contra effective effect. Although adverse events did not occur, most women in this study experienced side-effects from the oestradiol tablets. Frequent reported side-effects were reported painful swollen breasts, mood swings, headache and weight gain. This is important to include in counselling future patients. Regardless of the reported side-effects, most women stayed neutral or were (very) satisfied with the treatment. Results of this study support the hypothesis that ongoing bleeding is due to atrophy of the endometrium and form the basis to perform future research. The strong feature of this study is that it shows the first results of a treatment in long-term LNG-IUS users with irregular bleeding disturbances.

The present study has also some limitations. The sample size of this prospective, non-randomized cohort study is small (n=19). Hence, selection bias could be introduced which may limit our generalizability and limits also the ability for statistical analysis. There is also the risk of recall bias as women were asked for number of bleeding days in the last months (prior to starting oestradiol treatment). Finally, follow up was limited to three months. Long term follow-up is needed in future studies.

In conclusion, oestradiol may be a promising and safe treatment for irregular bleeding disturbances in long-term LNG-IUS users. First, we will start a large prospective study followed by a placebo-controlled trial evaluating addition of oestradiol as the treatment for irregular bleeding disturbances during long term LNG-IUS use.

Implications for practice and/or policy

Results of this feasibility study form the basis to perform future research.

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| Number of bleeding days* | Before oestradiol treatment (N=15) | After oestradiol treatment (N=16) | P value |
|--------------------------|-----------------------------------|----------------------------------|---------|
| No menstruation          | 0 (0)                             | 1 (0.5)                          |         |
| Normal menstruation      | 0 (0)                             | 8 (42)                           |         |
| Heavy menstrual bleeding | 2 (11)                            | 0 (0)                            |         |
| Intermensural bleeding   | 2 (11)                            | 3 (16)                           |         |
| Contact bleeding         | 3 (16)                            | 5 (26)                           |         |
| Spotting                 | 2 (11)                            | 2 (11)                           |         |
| Blood loss corresponding | 13 (68)                           | 6 (32)                           |         |
| section-niche            | 0 (0)                             | 0 (0)                            |         |

Table 3. At 3-month follow-up. a: Data are given as median (Interquartile Range), b: Wilcoxon signed rank test

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