Comparing the Effects of Continuous Hormone Replacement Therapy and Tibolone on the Genital Tract of Menopausal Women; A Randomized Controlled Trial

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Abstract

Introduction: Many postmenopausal women who are on hormone replacement therapy discontinue medications due to vaginal bleeding. Tibolone, a synthetic steroid, has minimal stimulatory effect on the endometrium. The aim of this study was to assess the effects of continuous HRT regimen and tibolone on the onset of vaginal bleeding and vaginal maturation value.

Materials and Methods: A total of 150 healthy women in postmenopausal period were randomly enrolled in this controlled clinical trial. Patients were randomly allocated into three groups, and were followed for six months. The first 50 women received 2.5 mg tibolone plus a Cal+D tablet (500 mg Calcium and 200 IU vitamin D) daily, the second 50 women received 0.625 mg conjugated equine estrogen and 2.5 mg medroxyprogesterone acetate (CEE/MPA) plus one Cal+D tablet daily, and the remaining 50 received only one Cal+D tablet per day and served as the control group. Symptoms were recorded using a questionnaire that assessed vaginal bleeding or spotting, vaginal dryness and intention to continue the medications. Vaginal maturation value was assessed by examining vaginal smears before and after the treatment. The results for the three groups were analyzed using statistical methods.

Results: In comparison with the control group, CEE/MPA and tibolone increased vaginal maturation value and decreased the frequency of vaginal dryness (p < 0.01). Women in tibolone group were more likely to continue the treatment regimen than those in the CEE/MPA or the control groups (p < 0.01).

Conclusion: Tibolone can serve as an appropriate choice for HRT as it has low rates of vaginal bleeding/ spotting episodes and high acceptance rate in postmenopausal women.

Keywords: Calcium, Hormone replacement therapy, Menopause, Tibolone, Vaginal bleeding, Vaginal maturation value.

Introduction

Many postmenopausal women use long-term hormone replacement therapy (HRT) and for most of them the potential benefits greatly outweigh its possible disadvantages or risks. Irregular bleeding occurs in up to 60% of HRT recipients which leads to discontinuation of the therapy in up to one in three users (1 - 3). The management of HRT-associated bleeding problems is invariably unsatisfactory, since there are no established methods for regulating or reducing the bleeding. This method is of clinical importance for a number of reasons. First, the potential...
health advantages of HRT, apart from the menopausal symptom relief, require longer use to be of benefit, although expectation of unscheduled vaginal bleeding may deter many women from starting HRT. Second, erratic bleeding in postmenopausal women may be a presenting symptom of malignancies of the cervix, endometrium or ovaries; therefore, current management protocols which involve invasive and costly investigations, may further result in anxiety.

Tibolone is a synthetic steroid with tissue-specific actions. Following rapid conversion to $3\alpha$- and $3\beta$-hydroxy-metabolites and the $\Delta4$-isomer, tibolone may engage the ER-$\alpha$, expressing estrogenic actions or the progesterone and androgen receptors and act as a progestogen-androgen (4). Tibolone has been administered in Europe as an effective alternative to continuous HRT in treating climacteric symptoms and vaginal atrophy, in improving libido and in preventing bone loss (5) with minimal stimulatory effect on the breasts (6, 7) or endometrium (8). These data, however, mostly result from observational studies, while results from randomized controlled trials are highly needed to reach a common conclusion.

This randomized controlled trial was undertaken to assess the effects of continuous HRT and tibolone regimen on the frequency of spotting/bleeding episodes and vaginal maturation value (VMV) in menopausal women.

Materials and Methods

A total of 150 healthy postmenopausal women, aged 45-60 years, whose last menstrual period was more than a year ago and had a plasma $17\beta$-estradiol $< 35 \text{ pg/ml}$ were enrolled in the study. The subjects were referred from Fadjr Hospital and a private clinic in Tehran, when they had attended to seek medical advice about their menopause. None of the participants had been treated with hormones known to influence vaginal bleeding during the preceding six months before the study. The initial screening included taking a medical history, and doing physical and gynecological examinations.

Patients were randomly allocated into three groups, according to a computer-generated list of random numbers, and were followed for six months. Fifty women received 2.5 mg tibolone (Tibofem, Cipla Ltd., India) plus one Cal-D tablet (Calcium 500 mg and vitamin D 200 IU), (Pharmachemi Ltd., Iran) on daily bases, 50 women received 0.625 mg conjugated equine estrogen plus 2.5 mg medroxyprogesterone acetate (CEE/MPA), (Aboreihan Ltd., Iran) plus a daily Cal+D tablet, and the remaining 50 women received only a Cal+D tablet and served as the control group.

Symptoms were recorded using a questionnaire that assessed vaginal bleeding or spotting, vaginal dryness and intention to continue the medications. The data were collected before and three and six months after the intervention. The following definitions were used for bleeding/spotting recordings:

Bleeding: any vaginal flow requiring more than one sanitary napkin per day.
Spotting: any vaginal flow requiring not more than one sanitary napkin per day.

Estrogenisation of the vagina was assessed by performing vaginal smears before and after the treatment by drawing the rounded part of a spatula from the upper portion of each lateral vaginal wall to the lower part of the organ. The resulting specimens were thinly smeared onto a slide and immediately fixed by 95% ethyl alcohol. Vaginal smears were examined by a skilled pathologist. Vaginal Maturation Value (VMV) was calculated by multiplying the percentage of cell types by a weighting factor and then adding up the total. The weights applied were: superficial cells 1.0; intermediate cells 0.5; Para basal cells 0.

Blood chemistry tests were performed for sex hormone binding globulin (SHBG) and the free estradiol index (FEI = estradiol $\times 100$/ SHBG). The levels of SHBG and estradiol were determined using ELISA (manufactured by IBL and DRG kits, respectively). Samples were handled in identical, blinded fashion throughout the study. They were analyzed in triplicate and in random order so as to reduce system bias and interassay variations. All the measurements were repeated after six months of commencing the study.

For a better evaluation of vaginal bleeding, the endometrium was assessed by sonography, and when endometrial double-wall thickness was more than 5 mm, additional biopsies were performed.

This study was approved by the Ethic Commit-
Statistical Analysis: The χ², Kruskal Wallis and Wilcoxon signed rank tests were used for the analysis of non-parametrical data. For the analysis of parametric data, Tukey adjustment for the comparison of means (ANOVA) and paired T-test were used. The significance level was set at $p < 0.05$. All confidence intervals were calculated at the 95% level.

Results

One-hundred and fifty postmenopausal women participated in this study. Four women withdrew from the study in the first month due to breast tenderness in three women and fear of breast cancer in another. During the study two women in the tibolone group, six women in the continuous HRT group and one in the control group interrupted the treatment before the third month due to vaginal bleeding or prolonged vaginal spotting. Finally, 47 women in the tibolone group, 42 women in the CEE/MPA group, and 48 women in the control group completed the six-month period of the study.

No individual with bleeding or spotting was found to have endometrial hyperplasia in endometrial biopsy.

There were no statistically significant differences among the three groups for parameters including age, age at menopause, gravidity, BMI and VMV shown in Table 1 before the treatment. The CEE/MPA group had higher incidence of bleeding and spotting episodes than the tibolone and control groups during the first three months (Table 2).

Table 1. Comparing the demographic data among the three groups of menopausal women on HRT, Tibolone and placebo (M ± SD or n (%))

|                    | Tibolone (n=49) | CEE/MPA (n=47) | Control (n=49) | P value |
|--------------------|-----------------|----------------|---------------|---------|
| Age (years)*       | 51.57 ± 3.18    | 51.10 ± 4.14   | 52.56 ± 4.16  | N.S     |
| Age at menopause (years)* | 49.07 ± 2.92   | 48.29 ± 2.91   | 48.65 ± 3.76  | N.S     |
| Gravidity**        |                 |                |               |         |
| <3                 | 10 (20.40)      | 9 (19.15)      | 9 (18.37)     | N.S     |
| 3-5                | 39 (79.60)      | 38 (80.85)     | 40 (81.63)    | N.S     |
| BMI (kg/m²)*       | 28.08 ± 3.30    | 29.14 ± 3.44   | 28.64 ± 3.51  | N.S     |
| VMV***             | 26.95 ± 18.52   | 33.14 ± 16.18  | 27.51 ±18.98  | N.S     |

*ANOVA test, **χ² test, ***Kruskal wallis test

Table 2. Bleeding / spotting (B/S) episodes among the three groups of menopausal women on HRT, Tibolone and placebo (M ± SD or n (%))

|                          | Tibolone group (n:49) | CEE/MPA group (n:47) | Control group (n:49) | Pa*       | Pb*       | Pc*       |
|--------------------------|-----------------------|----------------------|----------------------|-----------|-----------|-----------|
| Women reporting B/S episode during the 6 months | 14 (28.57) | 28 (59.54) | 9 (18.36) | <0.05 | N.S | <0.01 |
| Number of B episode during the first 3 months | 0.14 ± 0.45 | 0.69 ± 1.02 | 0.24 ± 1.30 | 0.03 | N.S | <0.01 |
| Number of B episode during the second 3 months | 0.12 ± 0.38 | 0.26 ± 1.09 | 0.18 ±1.27 | N.S | N.S | N.S |
| p**                      | N.S                  | <0.01                | N.S                  | -         | -         | -         |
| Number of S episode during the first 3 months | 0.20 ± 0.81 | 0.74 ± 1.21 | 0.22 ± 1.29 | <0.01 | N.S | <0.01 |
| Number of S episode during the second 3 months | 0.46 ± 1.21 | 0.36 ± 1.10 | 0.24 ± 1.33 | N.S | N.S | N.S |
| p**                      | N.S                  | <0.01                | N.S                  | -         | -         | -         |

*Pa, P-value between tibolone and CEE/MPA groups, Pb, P-value between tibolone and control groups, Pc, P-value between CEE/MPA and control groups, χ² test, Kruskal wallis test, Wilcoxon signed rank test
There were significant differences between the tibolone and CEE/MPA groups for SHBG, FEI and the number of women with vaginal dryness ($p < 0.001$). In comparison with the control group, CEE/MPA and tibolone increased VMV and decreased the number of women with vaginal dryness ($p < 0.01$). Women in the tibolone group were more willing to continue the medication in comparison with the CEE/MPA and the control groups ($p < 0.01$).

**Discussion**

The purpose of administering continuous HRT is to prevent scheduled bleeding and improve compliance among women wishing to remain amenorrheic. However, women on continuous HRT may still develop unscheduled bleeding, a serious and disturbing adverse event which may influence their decision to terminate or adhere to the treatment.

Accepted treatment modalities have been re-evaluated in varying estrogen-progestin dose ratios and alternative therapies have been considered in an attempt to alleviate their side-effects (9). Furthermore, investigators have focused on identifying the biological mechanisms responsible for the occurrence of unscheduled bleedings. Unscheduled bleedings may be influenced by factors such as menopausal age, BMI, the level of endogenous estrogens and its corresponding effect on mucosal changes in the genital tract (2).

The present study was designed to examine the effects of tibolone on the genital tract in postmenopausal women compared with that of continues HRT over a 6-month period.

In the present study, the prevalence of bleeding / spotting episodes in continues HRT was significantly higher as compared to tibolone and the controls. This study supports the results of studies that depict a lower bleeding/spotting episodes for tibolone intake, than continues HRT (2, 10, 11).

In order to acquire objective data, we practically used the hormonal milieu of the body and measured VMI. Vaginal maturation, as assessed by VMI, vaginal dryness and FEI are highly reliable predictors of genital tract mucosal changes following different treatment regimens. Significant differences in mean VMI, the number of women with vaginal dryness and FEI were observed in women in HRT and tibolone groups in comparison with the control group as similarly reported by several other investigators (12, 13).

We have shown here by cytological analysis that six months of tibolone therapy produces a significant, sustained oestrogenic effect on the vagina and tibolone significantly had lower bleeding/spotting episode rate. Tibolone decreased vaginal dryness, as seen in our previous study and improved sexual dysfunction in comparison with CEE/MPA (14).

As sexuality is an important aspect of health that affects the overall well-being of postmenopausal women, tibolone can be an appropriate choice for HRT in postmenopausal women as it has low bleeding/spotting episodes.

Tibolone increased fat free mass without any increase in weight or fat mass in postmenopausal (15). As adipose tissue has adverse effects on vaginal bleeding due to undesirable hormonal changes, this beneficial effect may cause low vaginal bleeding in individuals taking tibolone.

The limitation of this study was lack of endometrial assessment by regular sonography, unless vaginal bleeding or spotting was the case. In osteoporosis prevention and arterial effects of tibolone study (OPAL), endometrial thickness

**Table 3. Comparing the three groups of menopausal women before and after treatment with HRT, Tibolone and placebo (M ± SD or n (%))**

| Tibolone group (n=47) | CEE/MPA group (n=42) | Control group (n=48) | $P_a$** | $P_b$** | $P_c$** |
|----------------------|----------------------|----------------------|---------|---------|---------|
| **before** | **after** | **P** | **before** | **after** | **P** | **before** | **after** | **P** |
| VMV | 26.95±18.52 | 53.07±19.50 | <0.001 | 33.14±16.18 | 51.65±21.29 | <0.001 | 27.51±18.98 | 29.41±18.75 | N.S | N.S | <0.01 | <0.01 |
| Women who like to continue | - | 38(81.06) | - | 31(73.80) | - | 34(70.8) | - | 0.001 | 0.002 | N.S |
| Women with vaginal dryness | 36(73.46) | 33(73.46) | <0.01 | 34(72.34) | 20(40.00) | <0.001 | 36(72.00) | 37(74.00) | - | <0.001 | <0.01 | <0.001 |
| SHBG | 38.33±13.95 | 23.01±10.50 | <0.001 | 45.92±15.30 | 74.86±36.30 | <0.001 | 45.27±22.10 | 40.50±20.81 | N.S | <0.001 | <0.001 | <0.001 |
| FEI | 0.31±0.45 | 0.49±0.44 | 0.021 | 0.20±0.09 | 0.16±0.20 | N.S | 0.21±0.18 | 0.26±0.25 | N.S | <0.001 | 0.01 | <0.01 |

$P_a$, P-value between tibolone and CEE/MPA groups, $P_b$, P-value between tibolone and control groups, $P_c$, P-value between CEE/MPA and control groups,* paired t test, $\chi^2$ test, Wilcoxon signed rank test,** ANOVA (Tukey) test,kruskal wallis, $\chi^2$ test
measured by transvaginal ultrasound increased slightly during the first year on tibolone and CEE/MPA, but no further progression occurred after three years (11).

**Conclusion**

The rate of bleeding/spotting episodes decreased progressively in CEE/MPA group after three months of treatment. Tibolone was generally well-tolerated and the overall continuation rate was significantly higher. It seems that tibolone can be an appropriate choice for HRT due to lower rates of bleeding/spotting episodes and higher acceptance rate in postmenopausal women.

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