INDIVIDUALIZATION OF CUSTOM COMPOUNDED HORMONE THERAPY IN A PATIENT WITH CHEMOTHERAPY INDUCED PREMATURE OVARIAN INSUFFICIENCY AND IMPAIRED LIVER FUNCTION – CASE REPORT

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SUMMARY – Although the use of commercially manufactured hormone therapy (HT) to treat menopausal symptoms has declined during the past 12 years, the use of custom compounded HT seems to have increased. A 39-year-old woman with refractory anemia sustained premature ovarian insufficiency following allogeneic stem cell transplantation. After systemic biologic treatment (azacitidine) and corticosteroid therapy, besides extreme climacteric symptoms (Green Climacteric Scale, 59) and impaired quality of life, she also had elevated liver enzymes. Therefore, she was not a candidate for oral HT. Treatment was started with 17-beta estradiol patch 0.5 mg (Climara) together with micronized progesterone intravaginally, 2x100 mg (Utrogestan) for 3 months. She was not satisfied, so the custom compound HT started with 17-beta estradiol 0.5 mg gel 2x/day and micronized progesterone in liposomal gel 100 mg/daily. She was much better but she complained of low libido, decreased sex drive and emotional instability, so 1% testosterone gel was added. Now she was completely satisfied, Green Climacteric Scale was 8 and liver enzymes were normal. In conclusion, custom compound HT has the possibility of tailoring and adjusting therapy to the individual need, which has been the everlasting goal in menopause medicine and should be a good option for special clinical cases.

Key words: Custom compounded hormone therapy; Individualization; Premature ovarian insufficiency

Introduction

After the Women’s Health Initiative (WHI) study was published in the literature1, the use of hormone therapy (HT) has decreased dramatically. In 2010, the prevalence of HT use was only 4.7% in women over 40 years in the US as compared with 22.4% in 20002,3. Although data in the later WHI publication after 13-year follow-up differed from the initial reports and did not show an increase in cardiovascular risk in any age group (with the exception of venous thrombosis), the majority of women still refused to use HT4. However, individualization is the key approach including continuous ongoing assessment and use of new diagnostic tools for efficacy and risk assessment, and therapeutic adjustments as necessary; all these will ensure the best welfare of postmenopausal women5.

Custom compounded HT could be a form of personalized medicine whereby the dose, regimen and dosage forms are customized and based on the patient...
symptoms, hormone levels and personal preferences\(^a\). Since the Food and Drug Administration (FDA) approved HT, it has become the gold standard for therapeutic purposes; however, there are some special cases where FDA HT products do not fulfill individual patient demands. Therefore, in our case report, we present a patient with premature ovarian insufficiency (POI) because of refractory anemia and allogeneic stem cell transplantation, which seriously impaired her quality of life (QoL).

**Case Report**

A 39-year-old woman presented with a one-year history of amenorrhea. She had regular menstrual cycles from the age of 14 until after the allogeneic stem cell transplant for refractory anemia with excess blasts in September 2012. The post transplant period was complicated with chronic graft versus host disease (GvHD) of the skin and mucosa. She was treated with a combination of sirolimus and corticosteroids, which both were discontinued in July 2013, i.e. two months before examination at our institution. She continued taking therapy with mycophenolate until December 2014 when GvHD remission occurred and therapy was stopped. She remained in disease remission for 3.5 years after the transplant.

At presentation, she had normal body weight with body mass index of 21. She had normal secondary sexual characteristics, no signs of androgenism and no galactorrhea. She had very severe hot flushing and sweating at night, was unable to sleep and found extreme difficulty in concentrating. Her libido was significantly decreased. Her DEXA T-score L1-L4 was -1.6 SD, left femoral neck -0.2, total left hip -0.1 SD. She had no prior fragility fracture. Hormonal work-up confirmed elevated follicle-stimulating hormone, normal thyroid-stimulating hormone, normal prolactin, and normal basal and adrenocorticotropic hormone (ACTH) stimulated cortisol levels. Pelvic ultrasound examination showed small uterus with atrophic endometrium and involutive ovaries. Antimüllerian hormone or antral follicle count were not assessed since the patient declined to be referred to the Reproductive Unit, Department of Obstetrics and Gynecology, University Medical Centre Ljubljana, where these biomarkers are routinely assessed because she was not interested in evaluation of her ovarian reserve. We concluded that previous chemotherapy with multiple agents, including high dose busulfan and cyclophosphamide for stem cell transplant conditioning had led to premature ovarian failure causing her severe climacteric problems. Unfortunately, her liver enzymes were elevated in the post transplant period. Derangements of liver enzyme tests are often seen and usually attributed to drug interactions after allogeneic stem cell transplant. In addition, GvHD is a complication occurring in about half of these patients and in its acute form can affect the skin, liver and gut but is commonly seen as an overlapping syndrome affecting several organs. Due to elevated liver enzymes, the patient could not take oral HT, therefore another non-oral option should have been administered to improve her impaired QoL. The climacteric score sheet based on the Greene Climacteric Scale was obtained to objectify her complaints (Table 1). The hormones and liver enzyme laboratory tests before HT are reported in Table 2.

At first, the following FDA approved HT was administered: 17-beta estradiol in patch 50 mcg (Climara) once weekly + micronized progesterone 200 mg/day intravaginally (Utrogestan). After three months of treatment, the Greene Climacteric Scale of climacteric symptoms was repeated. There was no change in terms of QoL improvement related to the severity of climacteric symptoms. The patient’s score was 13 for anxiety, 10 for depression, 3 for sexual, 23 for psychological, 2 for physical, and 4 for vasomotor complaints. She was not satisfied with estrogen patch, she had allergic reactions around the patch, and complained of severe headache especially the first 2 days after the patch had been pasted. Therefore, the route of HT administration was switched to custom compounded HT (CC HT).

The 17-beta estradiol alcoholic gel 0.05% (50 mcg/g) 2x daily on the forearm + 10% (100 mg/g) micronized progesterone liposomal gel 2x daily on the chest was administered. After three months, the patient felt much better, with no vasomotor symptoms, but still had emotional instability and no libido. So the local 2% testosterone thick non-alcoholic gel 1x daily on the clitoris and labia for 2 months (to induce libido) was added, and then switched to systemic 1% testosterone micro in liposomal gel. After 3 months of this treatment, the patient achieved complete satisfaction. The Greene Climacteric Scale symptoms are shown in Table 3.

Three months after the CC HT was started, the hormone and liver enzyme laboratory tests were re-
peated (Table 4). We observed normalization of alkaline phosphatase, AST and ALT, and a significant decrease in gamma-glutamyl transferase, which were attributable to the successful GvHD treatment and immunosuppressive treatment cessation.

**Discussion**

Premature ovarian insufficiency is very common in women following allogeneic stem cell transplant. Ovaries are more susceptible with higher doses of chemotherapeutic agents, longer duration of treatment, and use of multiple agents. Artificial premature ovarian failure seemed to be abrupt cessation of reproductive hormonal function which led to impaired QoL. Therefore, HT was extremely important not only to improve her QoL but also to resume her normal life. Usually, for women with POI, a higher dose of estrogen is needed and maintained until the age of natural menopause.

Hormone therapy was the gold standard to enable women's life back, so tailoring the dose according to

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**Table 1. Green Climacteric Scale score sheet – before hormone therapy**

| Anxiety | Depression | Sexual | Psychological | Physical | Vasomotor |
|---------|------------|--------|---------------|----------|-----------|
| Average | >10        | >10    | 7             | 3        | 2         |
| Woman   |            |        |               |          |           |
| Average |            |        |               |          | 2         |
| Menopausal | >10 | >10 | 12            | 6        | 4         |
| Woman   |            | 3      | 24            | 2        | 6         |

Score: 14 10 3 24 2 6

The severity of problem is scored as follows: score 0 = none; 1 = mild; 2 = moderate; 3 = severe

**Table 2. Sexual hormones and liver enzymes before hormone therapy**

| Sexual hormones before HT | Liver enzymes before HT |
|---------------------------|-------------------------|
| S-estradiol, 0.081 nmo/L (0-0.11) | S-alkaline phosphatase, 3.25 μkat/L (1.74)* |
| S-progesterone, 0.64 nmol/L (0.2-3.2) | S-AST, 1.36 μkat/L (0.52)* |
| S-testosterone, 0.1 nmol/L (0.3-2.0)* | S-ALT, 2.82 μkat/L (0.56)* |
| S-DHEAS, 0.9 nmol/L (3.6-12.9)* | S-γ-GT, 7.49 μkat/L (0.63)* |
| S-FSH, 92 nmol/L* |                           |
| S-LH, 31 nmol/L* |                           |
| S-cortisol (basal), 238 nmol/L (101-536) | ACTH stimulated cortisol, 632 nmol/L |

* out of the normal range; S = serum; DHEAS = dehydroepiandrosterone sulfate; FSH = follicle-stimulating hormone; LH = luteinizing hormone; ACTH = adrenocorticotropic hormone; AST = aspartate aminotransferase; ALT = alanine aminotransferase; γ-GT = gamma-glutamyl transferase
the individual purpose was extremely important. The FDA approved HT satisfied most of the purposes for postmenopausal women, unfortunately, there were some exceptions.

The new International Menopause Society (IMS) recommendations for women’s midlife health and hormone therapy do not recommend using so-called ‘bio-identical hormones’ in the form of CC HT. Nevertheless, in our case, there were some objective obstacles that forced us to prescribe CC HT. Firstly, the woman had impaired liver function. Therefore, oral HT use was contraindicated concerning ‘the first pass through the liver’. Secondly, in Slovenia, there was a lack of transdermal FDA HT due to poor interest in HT use. Only one matrix patch system was available (17-beta estradiol, Climara 50 mcg). There was no gel or cream, either for estrogen or progesterone use. Moreover, transdermal testosterone therapy available in the dose appropriate for the woman was not FDA approved; the testosterone cream was available only in Australia.

Many CC HT prescribers worldwide have mixed estrogens in the form of Bi-Est (estradiol + estrone) or Tri-Est (estradiol + estrone + estriol), which is unnecessary either in terms of hormone metabolism or post-menopausal hormone levels. Estradiol is the main hor-

Table 3. Green Climacteric Scale after the custom compounded transdermal hormone therapy use

| Heart beat quickly and strongly | Feeling dizzy or faint | Feeling tense or nervous | Pressure or tightness in head or body |
|--------------------------------|-----------------------|------------------------|---------------------------------------|
| Feeling in sleeping            | Parts of body feeling numb or tingling |
| Excitable                      | Headaches             |
| Attacks of panic               | Muscle or joint pains |
| Difficulty in concentrating    | Loss of feeling in hands or feet |
| Feeling tired or lacking in energy | Breathing difficulties |
| Loss of interest in most things | Hot flushes |
| Feeling unhappy or depressed   | Sweating at night |
| Crying spells                  | Loss of interest in sex |

The severity of problem is scored as follows: score 0 = none; 1 = mild; 2 = moderate; 3 = severe

Table 4. Sex hormones and liver enzymes 3 months after custom compounded transdermal hormone therapy (CC HT) use

| Sex hormones after 3 months of CC HT | Liver enzymes after 3 months of CC HT |
|--------------------------------------|---------------------------------------|
| S-progesterone. 3.08 nmol/L (-3.2 nmol/L) | S-alkaline phosphatase. 1.66 |
| S-estradiol. 0.33 nmol/L (0.01-11.1 nmol/L) | S-AST, 0.37 |
| S-SHBG, 31 nmol/L (26.1-110.0 nmol/L) | S-ALT, 0.37 |
| S-testosterone (free). 16.1 nmol/L (3.5-20.9 nmol/L) | S-γ-GT, 2.13* |
| S-FSH, 42.7 nmol/L | |
| S-LH, 23.7 nmol/L | |

* out of the normal range; S = serum; SHBG = sex hormone binding globulin; FSH = follicle-stimulating hormone; LH = luteinizing hormone; AST = aspartate aminotransferase; ALT = alanine aminotransferase; γ-GT = gamma-glutamyl transferase
mone produced by the ovary that is decreased significantly in postmenopause; estradiol is a hormone that remains almost at the same premenopausal level due to extra-genital production from the androgen[11-14]. So, a mixture of all estrogens should not improve QoL in postmenopausal women better than estradiol alone.

Salivary or serum hormone testing is not necessary, especially in order to adjust the dose of the hormone regarding hormone levels in postmenopausal women. There are at least two reasons why hormone testing is not appropriate for adjusting the dose of the hormones: first, estrogen, as well as progesterone or testosterone, have a pulsatile hormone release, so the concentration in serum or saliva changes several times a day; and second, hormone release is connected with dietary patterns and interactions with other medications, so the hormone levels in serum or saliva need not necessarily reflect the hormone levels in the target organ[15,16]. Hormone testing is reasonable in women with POI to examine the basal pretreatment hormone levels, and repeat testing after 3 months of therapy to follow-up the body response to hormone treatment according to the route of administration. Thus, hormone testing should not be the basis for adjusting the HT dose, but rather an opportunity for patient follow-up concerning the prescribed dose and route of administration.

Recent studies confirmed that the risk of venous thromboembolism, as well as its impact on fibrinolytic and coagulation parameters varied according to HT formulation[17-21]. The greatest were recorded in the users of oral estrogen-progestin HT. Transdermal route of administration seems to be much safer, with less side effects concerning thrombotic risk[22].

Using transdermal micronized progesterone was an issue of great concern regarding endometrial safety. The effectiveness of these preparations in protecting the endometrium is controversial due to the very low serum levels that are achieved. A recent publication has finally confirmed that topical alcohol-based gels appear to yield luteal-phase serum progesterone levels which satisfactorily protect the endometrium[21]. We used liposomal gel, which ensures high bioavailability and deep penetration of progesterone, even better than alcohol-based gel[21,25]. The 10% progesterone liposomal gel twice daily enables 200 mg micronized progesterone, which ensures safe endometrial protection.

In conclusion, our tasks as practitioners should be the responsibility to achieve the best QoL in our patients with special circumstances, using either FDA approved HT with natural hormones, or CCHT, fulfilling the criteria of individualization and tailoring therapy upon personal needs. Therefore, we could achieve the criteria of good clinical practice in menopausal medicine.

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Sažetak

INDIVIDUALIZACIJA MAGISTRALNOG HORMONSKOG LIJEČENJA KOD BOLESNICE S KEMOTERAPIJOM INDUCIRANOM PRIJEVREMENOM INSUFICIJENCIJOM JAJNIKA I SMANJENOM JETRENOM FUNKCIJOM: PRIKAZ SLUČAJA

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Premda je upotreba komercijalno pripravljenih hormonskih terapije u liječenju klimakteričnih simptoma u posljednjih 12 godina u padu, čini se da je upotreba magistralnih hormonskih pripravaka u porastu. Žena u dobi od 39 godina s refraktornom anemijom doživjela je prijevremenu insuficijenciju jajnika nakon transplantacije matičnih stanica. Nakon sistemskog biološkog liječenja azacitidinom i kortikosteroidima, uz izravne klimakterične tegobe (Greenov indeks 59) i smanjenju kvaliteta života imala je povišene jetrene enzime. Zbog toga nije bila kandidat za oralnu hormonsku terapiju. Započeto je liječenje 17-beta estradiolom u obliku naljepka od 0,5 mg (Climara) zajedno s mikroniziranim progesteronom intravaginalno 2x100 mg (Utrogestan) kroz 3 mjeseca. Nije bila zadovoljna terapijom pa su joj propisani magistralni pripravci.

Započelo se s primjenom 17-beta estradiolom u obliku 0,5 mg gela 2x/dan i mikroniziranog progesterona u liposomalnom gelu 100 mg/dnevno. Bolje se je osjećala, ali još uvijek se žalila na smanjeni libido i emocionalnu nestabilnost pa je dodan 1% testosterona. Sad je bila potpuno zadovoljna terapijom, Greenova klimakterijska ljestvica bila je 8, a jetreni enzimi su se normalizirali. U zaključku, magistralni hormonski pripravci pružaju mogućnost titracije i prilagođavanja terapije individualnim potrebama, što je stalni cilj u menopauzalnoj medicini i mogao bi biti dobra mogućnost za posebne slučajeve.

Ključne riječi: Magistralno pripravljeni hormoni; Individualizacija; Prijevremena insuficijencija jajnika

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