GUIDELINES AND POSITION STATEMENTS

Peri- and postmenopause—diagnosis and interventions interdisciplinary S3 guideline of the association of the scientific medical societies in Germany (AWMF 015/062): short version

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Abstract
This short version of the interdisciplinary S3 guideline “Peri- and Postmenopause—Diagnosis and Interventions” is intended as a decision-making instrument for physicians who counsel peri- and postmenopausal women. It is designed to assist daily practice. The present short version summarizes the full version of the guideline which contains detailed information on guideline methodology, particularly regarding the critical appraisal of the evidence and the assignment of evidence levels. The statements and recommendations of the full version of the guideline are quoted completely in the present short version including levels of evidence (LoE) and grades of recommendation. The classification system developed by the Centre for Evidence-based Medicine in Oxford was used in this guideline.

Keywords Perimenopause · Postmenopause · Hormone replacement therapy · Diagnosis · Treatment · Risk communication

Introduction
Peri- and postmenopausal women often seek medical assistance because of climacteric symptoms (e.g. vasomotoric symptoms). They want information (a) about the physiological changes during the menopausal transition period and (b) ways to alleviate their symptoms to improve their quality of life. During peri- and postmenopause, symptoms may change. Dysfunctions and diseases may develop that may or may not depend on the changes of sex hormones. Perimenopausal women often ask for information about how to prevent diseases that are typically associated with the perimenopausal transition and the postmenopause. The current S3 guideline, Peri- and Postmenopause—Diagnosis and Interventions’ is based on the previous interdisciplinary S3 guideline, Hormone therapy in perimenopause and postmenopause’ [1, Appendix]. The authors of the current S3 guideline intended to take a broader spectrum of issues of the peri- and postmenopause into account and provide evidence-based recommendations to counsel women in this period of life.

See “Appendix” for guideline group.

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Diagnosis and treatment

Evidence-based recommendation

| Level of Evidence | Grade of recommendation | Degree of consensus |
|-------------------|-------------------------|---------------------|
| LLA               | A                      | ++                  |

Peri- and Postmenopause in women aged 45 or older should be diagnosed based on clinical parameters.

FSH should be used to diagnose peri- and postmenopause only in women between 40 and 45 years with climacteric symptoms or in women under the age of 40 years with suspected indicators of premature ovarian insufficiency.

The different phases of the menopausal transition can be mainly diagnosed by clinical criteria. Hormone measurements are usually not necessary [1, 2]. The following terms should be used for healthy women older than 45 years who have climacteric symptoms without laboratory tests:

- Perimenopause—women with irregular menstrual cycles and possibly vasomotor symptoms.
- Postmenopause—women without a menstrual bleeding for at least 1 year in the absence of interventions which may cause amenorrhea (e.g. oral contraceptives).
- Peri-/postmenopause—women after hysterectomy with vasomotor symptoms. In women who are using hormone replacement therapy (HRT), it may be difficult to define the menopausal status.

In women older than 45 years, measurements of anti-Müllerian hormone, inhibin A, inhibin B, antral follicle count or ovarian volume should not be used to diagnose peri- or postmenopause. Follicle stimulating hormone (FSH) measurement may be used in women between 40 and 45 years with climacteric symptoms or irregular menstrual cycles and in women younger than 40 years with suspected premature ovarian insufficiency [1, 2].

Symptoms

Vasomotor symptoms such as hot flushes and night sweats are the most common symptoms of the climacteric syndrome in peri- and postmenopausal women. Other symptoms such as sleep disturbances, mood swings, anxiety, and sexual impairment may also be associated with the peri- and postmenopause. In contrast to the vasomotoric symptoms, it is not clear whether the other symptoms are caused by hormonal changes. Loss of estradiol is the main cause of the climacteric syndrome, but the exact role of serum and tissue level changes of estradiol and other hormones such as androgens and gonadotropins is unclear.

Frequency and duration of all climacteric symptoms vary considerably and may be influenced by a number of factors such as general health and well-being, socioeconomic status, cultural influence, and other factors. The prevalence of hot flushes has been reported to be between 14 and 51% among premenopausal women, about 50% among perimenopausal women, and between 30 and 80% among postmenopausal women [3]. The median duration of hot flushes is 7.4 years. Longer durations typically occur when climacteric symptoms already develop in the premenopause [4].

Information

Peri- and postmenopausal women should be informed about the phases of peri- and postmenopause, the diagnosis, typical symptoms (vasomotor symptoms, sleep disorders, mood changes, urogenital symptoms, sexual impairment, changes in lifestyle and other measures which may affect health and well-being). In addition, they should be informed about the benefits and risks of different treatments strategies and the long-term effects of peri- and postmenopause on female health. Peri- and postmenopausal women should also be informed about different treatment options such as hormonal interventions, non-hormonal interventions and non-pharmacologic interventions.

Therapeutical interventions

Evidence-based recommendation

| Level of Evidence | Grade of recommendation | Degree of consensus |
|-------------------|-------------------------|---------------------|
| 1a                | A                      | ++                  |

Women with vasomotor symptoms should be offered hormone replacement therapy (HRT) after information about short- (up to 5 years) and long-term risks and benefits. Women with an intact uterus may receive estradiol replacement therapy with an appropriate progestin component (EPT), women after hysterectomy may receive estradiol replacement therapy (ET) only.
Evidence-based recommendation

Level of Evidence 3 Grade of recommendation A Degree of consensus + +

Serotonin reuptake inhibitors (SSRIs), serotonin noradrenaline reuptake inhibitors (SNRIs), clonidine, and gabapentin should not be offered as treatment of first choice for the treatment of vasomotor symptoms.

Evidence-based recommendation

Level of Evidence 1b Grade of recommendation 0 Degree of consensus + +

Cognitive behavioural therapy (CBT), isoflavones, and cimicifuga preparations may be used to treat vasomotor symptoms.

* See “Appendix” for dissenting opinion of the GPT

Vasomotoric symptoms

Estradiol is the most efficient means of therapy of climacteric symptoms. Therefore, HRT should be offered all women with moderate to severe climacteric symptoms with significant impairment of their quality of life. For women with an intact uterus, EPT is the most efficient treatment of vasomotor symptoms. Hysterectomized women should only receive ET. The progestin component of EPT can be applied cyclically or continuously. Potent progestins such as synthetic progestins should be used for at least 10–12 days per month. Shorter treatment periods, or the use of natural progesterone and dydrogesterone instead of synthetic progestins may increase the risk of endometrial hyperplasia and endometrial cancer. One must consider that different HRT preparations or routes of application have different risk profiles. Transdermal application may have a better risk–benefit ratio and should be preferred over oral preparations. Women who desire phytotherapy should be informed that numerous preparations exist, and that the safety of these preparations is often unclear. In addition, combinations of different phytotherapy compounds may have different effects and may interact with concurrent medications [1, 2].

Changes of sexual function

Evidence-based recommendation

Level of Evidence 1b Grade of recommendation 0 Degree of consensus + +

Peri- and postmenopausal women with loss of libido can be offered testosterone treatment after psychosexual exploration and when HRT has not been efficient. They should be informed about off-label use. Peri- and postmenopausal women may observe changes in sexual function. They should be asked about signs of vaginal atrophy such as dyspareunia. Loss of libido is common in ageing women. Low levels of estrogen and testosterone may be associated with loss of libido. However, there is no strict association between sex hormone levels and libido as reasons for loss of libido are complex. A psychotherapeutic intervention may be beneficial. Treatment with testosterone may be considered when HRT was not efficient for improving libido. Moisturizers and lubricants may be used alone or in addition to vaginal estrogen preparations. All these interventions are usually sufficient to reduce vaginal symptoms such as dryness, pain on intercourse or vaginal discharge. When estrogens are used for the treatment of vaginal dryness, estriol (E3)-containing preparations should be preferred. E3 may be used efficiently in low or ultra-low doses. The dosage may be increased if symptoms are not relieved. When treatment is terminated, symptoms typically re-occur. Side effects of vaginal ET are rare [1, 2].

Urogenital atrophy

Evidence-based recommendation

Level of Evidence 1b Grade of recommendation A Degree of consensus +++

Women with symptomatic urogenital atrophy should be offered lubricants alone or together with a vaginal ET. The treatment duration should be individualized.

Starting and stopping HRT

After starting HRT, women should consult their gynaecologist regularly (initially after 3 months) to monitor treatment effectiveness and tolerability of the therapy. Women should be advised to consult their gynaecologist in case of atypical bleeding. Modification of HRT has to be considered for several reasons during different phases of peri- and postmenopause. When treatment is well tolerated and no pathological symptoms occur, yearly gynecologic consultations are appropriate. Vaginal ultrasound examinations and endometrial thickness measurements should not be used routinely during HRT [5].

If it is determined that HRT should be stopped, two options can be offered: gradually reducing HRT or ceasing immediately. Gradually reducing may limit recurrence of symptoms in the short term. Whichever method is chosen, it has no influence on the recurrence rate of symptoms in the longer term.
Urogynecology

Urinary incontinence

Evidence-based statement
Level of Evidence 1a Degree of consensus +++
Vaginal ET may improve urinary incontinence in postmenopausal women

Evidence-based recommendation
Level of Evidence 1a Grade of recommendation A Degree of consensus ++
Patients should be informed before systemic ET or EPT that these therapies may lead to occurrence or worsening of urinary incontinence

Evidence-based recommendation
Level of Evidence 1a Grade of recommendation A Degree of consensus ++
Postmenopausal women with urinary incontinence should be offered pelvic floor training and vaginal ET

Randomised controlled trials have shown that in women with urinary incontinence, vaginal ET leads to significant improvements compared to placebo. However, a significantly higher prevalence of incontinence was documented when ET was applied systemically. Specifically, prevalence of urinary incontinence was doubled compared to placebo. EPT also leads to increases of urinary incontinence. However, the effect was smaller than in ET users [6, 11].

Overactive bladder

Evidence-based statement
Level of Evidence 1b Degree of consensus +++
Systemic HRT may lead to worsening of urinary incontinence. Women with overactive bladder may be offered vaginal ET

Evidence-based recommendation
Level of Evidence 1b Grade of recommendation 0 Degree of consensus ++
After exclusion of urological diseases, women with symptoms of urgency and frequency may be offered local ET

Urinary tract infections

Evidence-based statement
Level of Evidence 2b Degree of consensus +
Changes in the vaginal milieu of postmenopausal women may predispose to urinary tract infection, especially in older women

Evidence-based recommendation
Level of Evidence 2a Grade of recommendation B Degree of consensus ++
Recurrent urinary tract infections in postmenopausal women should preferably be treated with vaginal ET instead of antibiotics

Two small studies have demonstrated that vaginal ET reduced the frequency of urinary tract infections. Oral ET did not lead to a reduction of recurrent urinary tract infections [6, 8, 12, 16].

Cardiovascular disease

Evidence-based recommendation
Level of Evidence 2b Grade of recommendation B Degree of consensus ++
The risk for cardiovascular disease in peri- and postmenopausal women varies depending on their risk profile. These risk factors should be controlled optimally to exclude contraindications for HRT. Therefore, cardiovascular risk factors have to be identified and treated before HRT is initiated

Because of the various limitations of the trials, mainly the WHI, statements and recommendations on the effects of HRT on cardiovascular diseases are based on a moderate level of evidence. Large randomized intervention trials have shown no general protection against cardiovascular diseases by HRT but indicate a neutral or even negative effect of HRT in postmenopausal women [17]. The analyses of data including recent studies indicate the need for caution, but individualized treatment may have little risk or even cardiovascular
benefit. The transdermal application has advantages over the oral form [18]. In any case, the prerequisite seems to be the control of the conventional risk factors for cardiovascular events such as heart attack and stroke. Regardless of the study results, previous vascular events may be contraindications for HRT [1, 2].

**Thromboembolism**

| Evidence-based recommendation |
|--------------------------------|
| Level of Evidence | Grade of recommendation | Degree of consensus |
| 2a | A | + + |

Women should be informed that risk of thromboembolism is higher during oral compared to transdermal ET and EPT.

The highest vascular risks of oral HRT are venous thrombosis and thromboembolism [1, 2, 19, 21]. ET and EPT double the risk by about two cases per 1000 women per year [22, 25]. Estrogens and progestogens have thrombogenic effects that are dose dependent and substance specific, which are more pronounced in the initial phase of therapy and increase with age, weight and genetic predisposition [20, 21]. With low-dose transdermal therapy, there was no evidence of an increased risk of thromboembolism probably due to the lack of a first-pass effect in the liver [19, 21].

**Cerebrovascular events**

| Evidence-based recommendation |
|--------------------------------|
| Level of Evidence | Grade of recommendation | Degree of consensus |
| 2b | A | + + |

Women should be informed that oral EPT, but not transdermal ET may increase the risk for ischemic cerebrovascular events. Absolute risk for cerebral stroke is very low in younger women.

Ischemic strokes are the second most common and because of possible persistent disabilities very serious risk of oral HRT [2, 18, 19, 26]. In absolute numbers, the risk increases by 1 case per 1000 women per year during oral ET or EPT intake [22, 27, 28]. An increase in the risk of stroke may be avoidable using transdermal HRT up to a dose of 50 µg [1, 2].

**Coronary heart disease**

| Evidence-based recommendation |
|--------------------------------|
| Level of Evidence | Grade of recommendation | Degree of consensus |
| 2b | A | + + |

Women should be informed that EPT does not increase their cardiovascular risk or has only a minor effect. ET does not increase cardiovascular risk or may even reduce it. HRT is not appropriate for the primary or secondary prevention of coronary heart disease. HRT should be started before the age of 60 years for the treatment of climacteric symptoms.

Neither ET nor EPT have significant influences on the risks for coronary heart disease in primary and secondary prevention [17, 22, 23, 25]. However, HRT initiated within the first 10 years after menopause appears to be associated with a lower coronary event rate, while starting 20 years or more after menopause significantly increases the risk [1, 2, 19, 25].

**Osteoporosis**

| Evidence-based statement |
|--------------------------|
| Level of Evidence | Degree of consensus |
| 1a | +++ |

HRT significantly reduces the risk for osteoporotic fractures.

**Prevention**

Excessive weight reduction leads to an increased risk for atraumatic fractures. A body mass index below 20 should be avoided. Consequently, weight gain is associated with reduced risk of atraumatic fractures. Weight gain above a body mass index of 35 should be avoided. Exercise and sufficient intake of calcium and vitamin D is recommended [1, 2, 29, 32].
Treatment

A number of therapies are licensed for the treatment of osteoporosis: estrogen (in combination with progestin in non-hysterectomised women), bisphosphonates, selective estrogen receptor modulators (SERMs), and a human monoclonal antibody against the RANK ligand (Denosumab) and parathyroid hormone (teriparatid). The efficiency of EPT or ET for primary prevention of osteoporotic factors has been demonstrated in observational studies and randomized controlled trials. The current version of the S3 Guideline of the German Association of Osteology (DVO) recommends that prevention or treatment of osteoporosis by HRT is feasible when women have concurrent climacteric symptoms or do not tolerate or have contraindications against licensed treatments for the prevention or treatment of osteoporosis [1, 2, 29, 32].

The reduction of fractures by HRT is independent of the duration of HRT and the age at treatment initiation. In addition, the risk reducing effect of HRT seems to persist at a lower level after its termination. Since osteoporosis is a chronic disease, treatment is usually performed for longer durations depending on the assumed baseline risk for fractures. The mean duration is 3–4 years. In accordance to the results of the WHI, the following risk reducing effects can be expected: − 23 fractures per 1000 HRT users (confidence interval [CI] 10–33), − 20 non-vertebral fractures per 1000 HRT users (CI 6–30), − 8 vertebral fractures per 1000 HRT users (CI − 18 to + 9), − 0 hip fractures per 1000 HRT users (CI − 10 to + 4), − 36 wrist fractures per 1000 HRT users (CI 19–43) [1, 2]. In women with premature ovarian insufficiency, HRT or oral contraceptives should be used until the age of natural menopause to prevent osteoporosis and osteoporotic fractures.

Dementia, depression and mood changes

| Evidence-based recommendation |
|-------------------------------|
| Level of Evidence | Grade of recommendation | Degree of consensus |
| LLA               | A                        | +++                  |

Peri- and postmenopausal women should be informed that it is unclear whether HRT before the age of 65 has an influence on the risk of dementia. Peri- and postmenopausal women should be informed that it is unclear whether HRT before the age of 65 has an influence on the risk of dementia.

HRT and cancer risk

**HRT and breast cancer risk**

| Evidence-based recommendation |
|-------------------------------|
| Level of Evidence | Grade of recommendation | Degree of consensus |
| 1a                | A                        | +++                  |

Women considering HRT should be informed that HRT (EPT/ET) may lead to a small or no increased risk of breast cancer. Breast cancer risk depends on HRT formulations and treatment duration and is reduced after the end of treatment.

EPT

EPT may increase breast cancer risk. This was shown in a number of meta-analyses that included observational studies as well as randomized controlled trials (RCTs). In the randomized Women’s Health Initiative (WHI) trial, EPT led to an increased relative risk of breast cancer of 1.26 with 8 additional cases per 10,000 women per year [22, 35]. Cohort studies have shown that continuous combined EPT leads to higher increases in breast cancer risk than sequential EPT.
RCTs and observational studies demonstrated that current users of EPT have an increased breast cancer risk. However, this risk is reduced after stopping HRT. When HRT was initiated at or around the age of menopause, breast cancer risk increases were higher than in women who initiated HRT more than 5 years after menopause. EPT-containing progesterone leads to smaller increases of breast cancer risk compared to EPT-containing synthetic progestins [36]. However, progesterone has a smaller anti-proliferative effect on the endometrium than synthetic progestins. In the WHI study, breast cancer mortality was increased in HRT users in the short term, but not after 18 years of follow-up [1, 2, 37].

**ET**

In contrast to a large number of observational studies, the WHI trial demonstrated a significant risk reduction for breast cancer in ET versus placebo users [38]. The mean duration of ET was 7 years. Three smaller RCTs did not find significant differences in breast cancer risk of women using ET compared to placebo [39, 41]. Observational studies have shown a small increase in breast cancer risk in women using ET. The duration of ET treatment leading to an elevated breast cancer risks is controversial. Breast cancer-specific mortality was reduced in ET uses in the 18-years follow-up in the WHI trial [37]. However, this calculation is based on a small number of breast cancer cases.

In summary, HRT may be associated with increased breast cancer risk. The elevation is relatively small and has to be included in the individual risk–benefit calculation before initiating HRT for climacteric symptoms.

**Vaginal ET and breast cancer risk**

Vaginal ET may lead to increases of systematic estrogen levels. It is not known whether it may also lead to increases of breast cancer risk. Ultra-low doses of vaginal ET (e.g. 0.03 mg estriol, 2–3 applications per week) lead to good clinical effects on vaginal symptoms. It is unlikely that such an ultra-low-dose vaginal ET has a causal influence on breast cancer risk even when the therapy is used chronically [42].

**HRT after breast cancer**

| Evidence-based recommendation |
|-----------------------------|
| Level of Evidence 2b | Grade of recommendation | Degree of consensus +++
| HRT should not be used in women after breast cancer. In selected cases it may be used after ineffective non-hormonal treatments and severe limitations of quality of life |

Meta-analyses including observational studies and RCTs could not demonstrate an increased risk of relapse of breast cancer after using HRT [43, 46]. However, these studies have considerable methodological limitations such as low numbers of cases and short follow-up. The HABITS trial, an RCT, demonstrated an increase of breast cancer relapse in breast cancer survivors using HRT [44, 45]. However, the number of only 442 women in this study was small. The Stockholm randomized trial with 378 breast cancer survivors found no increased risk of relapse in the HRT arm. Therefore, it is not possible to make reliable conclusions on the oncological safety of HRT among breast cancer survivors. In the randomised controlled LIBERATE trial 3000 women with vasomotoric symptoms after breast cancer were treated with tibolone or placebo. Tibolone led to increased risk of breast cancer (hazard ratio [HR] 1.40) after a median follow-up of 3.1 years [47].

**Vaginal ET after breast cancer**

Women who have been or are being treated for breast cancer may complain about vaginal symptoms such as dryness, dyspareunia, and pain. They use vaginal ET six times more often than systemic HRT. Vaginal ET may lead to increases in serum estrogen levels. Vaginal ETs are different in type, dosage, and frequency of application. In Germany, estriol is frequently used for vaginal ET. Ultra-low-dose estriol in combination with lactobacillus acidophilus was tested in a small study on its safety and efficacy. In this study, vaginal atrophy was significantly improved and serum levels of estriol were only increased during the initial period of ET. After 4 weeks of treatment, levels of estriol were slightly elevated in 50% of users while not at all in the remaining 50%. It can be concluded that ultra-low-dose vaginal estriol therapy (0.03 mg, 3 applications per week) may be used in breast cancer survivors if non-hormonal alternatives had insufficient results [48].
HRT and endometrial cancer

Evidence-based statement
Level of Evidence 2 Degree of consensus +++
HRT containing an estrogen without a progestin is a risk factor for endometrial cancer in non-hysterectomized women. The effect is dependent on the duration of treatment

Evidence-based statement
Level of Evidence 2 Degree of consensus + +
Continuous combined HRT with conjugated equine estrogens and medroxyprogesterone acetate for an average duration of 5.6 years reduced the risk of endometrial cancer

Evidence-based statement
Level of Evidence 2 Degree of consensus + +
Continuous combined HRT for less than 5 years does not increase risk of endometrial cancer

Evidence-based statement
Level of Evidence 3 Degree of consensus + +
Long-term treatment with of continuous combined HRT for more than 6 or 10 years may lead to an increased risk of endometrial cancer

Evidence-based statement
Level of Evidence 4 Degree of consensus +
The use of progesterone or dydrogesterone in continuous combined HRT may increase the risk of endometrial cancer

Evidence-based statement
Level of Evidence 3 Degree of consensus +++
Sequential combined HRT may increase the risk of endometrial cancer. The effect depends on duration, type and dose of progestin

Evidence-based statement
Level of Evidence 3 Grade of recommenda- tion A Degree of consensus ++++
Sequential combined HRT for a duration of less than 5 years including a synthetic progestin does not increase the risk of endometrial cancer risk

Vaginal ET and endometrial cancer risk

Clinical studies did not show increased rates of endometrial hyperplasia after vaginal ET. Therefore, it is not recommended to add a progestin to vaginal ET. However, there are no data on endometrial safety of vaginal ET when used for more than 1 year [5].
**HRT after endometrial cancer**

Evidence-based statement

Level of Evidence 2b  
Degree of consensus +++

The effect of HRT on the risk of relapse after treatment of endometrial cancer has not been investigated sufficiently

Evidence-based recommendation

Level of Evidence 2b  
Grade of recommendation EK  
Degree of consensus + +

In patients who have been treated for endometrial cancer, HRT may be used for the treatment of climacteric symptoms if they have severe quality of life limitations and in whom non-hormonal treatments were not effective.

A recent meta-analysis has included only one randomized trial and five observational studies assessing the safety of HRT in women who have been treated for endometrial cancer. Of note, the risk of endometrial cancer relapse was reduced in HRT users. However, most studies included in this meta-analysis were retrospective observational studies. Therefore, the evidence regarding the oncological safety of HRT among endometrial cancer survivors is limited. It can only be concluded that HRT after early stage endometrial cancer may not lead to a relevant increase of the risk of relapse [5, 57, 59].

**Vaginal ET after endometrial cancer**

Evidence-based recommendation

Level of Evidence 4  
Grade of recommendation A  
Degree of consensus + +

Patients with symptoms of vaginal atrophy after treatment of endometrial cancer should be treated primarily with moisturizers or lubricants.

Consensus-based recommendation

EK  
Degree of consensus + +

After treatment of endometrial cancer, vaginal ET may be used if lubricants or cremes have proven to be ineffective.

Patients who have been treated for endometrial cancer may complain vaginal dryness, pain, dyspareunia, vaginal bleeding, and urinary incontinence. This may lead to sexual dysfunction and reduction of quality of life. Since it cannot be excluded that vaginal ET may lead to increase the risk of relapse, endometrial cancer survivors should be treated with non-hormonal moisturizers or lubricants [60]. Ph-stabilizing preparations with a pH between 4 and 4.5 have been shown to be efficient in this indication [61]. Local ET may alleviate the symptoms of vaginal atrophy after radiotherapy of the vagina [62]. Retrospective observational studies did not show increased rates of relapse after vaginal ET [63]. However, the evidence is limited and does not prove oncological safety [5].

**HRT and ovarian cancer risk**

Evidence-based recommendation

Level of Evidence 2a  
Grade of recommendation A  
Degree of consensus + +

Women considering HRT should be informed that ET and EPT may increase ovarian cancer risk. This effect has been observed after treatment durations of less than 5 years and is reduced after stopping treatment.

The Collaborative Group on Epidemiological Studies of Ovarian Cancer performed a meta-analysis of 52 studies. This included data from 21,488 postmenopausal women with ovarian cancer. Data from prospective trials showed that HRT users had increased risk for ovarian cancer after HRT durations of less than 5 years (RR 1.43). The absolute increase of risk is 1 in 1000 after 5 years and 1 in 1600 women using HRT [64].

**HRT after ovarian cancer**

Evidence-based statement

Level of Evidence 2b  
Degree of consensus + +

The safety of HRT after treatment of ovarian cancer is unclear.

Evidence-based recommendation

Level of Evidence 2b  
Grade of recommendation 0  
Degree of consensus +++

HRT may be used after treatment of ovarian cancer and appropriate consultation with the patient.

After treatment, ovarian cancer patients may experience natural or therapy-induced menopause. These patients may suffer from climacteric symptoms. Younger women may develop estrogen-associated diseases such as coronary heart disease and osteoporosis [65, 70]. There are only
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few studies that examined the safety of HRT after ovarian cancer [71]. Three observational studies could not demonstrate increased risk of relapse [66, 68, 69]. However, there are a number of methodological limitations. A randomized study with 150 patients showed improved survival in the group of women treated with HRT after a median follow-up of 19.1 years (HR 0.63; 95% CI 0.44–0.90). The authors conclude that HRT after ovarian cancer is safe. However, this study has a number of limitations and the safety of HRT after ovarian cancer remains unclear [67]. Since none of the studies showed an increased, but rather a risk reduction, HRT may be considered when women complain of severe climacteric symptoms. Especially in younger women with iatrogenic menopause after ovarian cancer treatment, the risk of mortality due to coronary heart disease is increased. In these women ET should be considered.

**HRT and risk for colorectal cancer**

| Evidence-based recommendation |
|-----------------------------|
| Level of Evidence 2a | Grade of recommendation A | Degree of consensus +++ |
| Women should be informed that HRT may reduce the risk of colorectal cancer. HRT should not be used for colorectal cancer prevention |

Reproductive factors may influence the risk of colorectal cancer. Studies have shown that HRT has no effect or reduces the risk for colorectal cancer [72, 73]. In the WHI trial, the risk of colon cancer was reduced by 37% in EPT users. ET did not have any effect on colorectal cancer risk [74]. Observational studies had conflicting results. The majority of these studies showed a risk reduction after HRT use. A meta-analysis including data from RCTs and observational studies showed that EPT as well as ET significantly reduced the risk of colorectal cancer (RR 0.74; 95% CI 0.68–0.81 and RR 0.79; 95% CI 0.69–0.91, respectively). A recent publication of data from five Danish cancer registries including 1.1 million women also demonstrated that ET and EPT reduced the risk of colorectal cancer with a minimal effect of EPT on rectal cancer [73].

**Premature ovarian insufficiency (POI)**

| Evidence-based recommendation |
|-----------------------------|
| Level of Evidence 2b | Grade of recommendation B | Degree of consensus +++ |
| Women with POI should be informed about the importance of HRT or combined oral contraceptives at least until the age of natural menopause if there are no contraindications against HRT or combined oral contraceptives |

Premature ovarian insufficiency (POI) or premature ovarian failure (POF) is defined as loss of ovarian function before the age of 40 and affects 1% of women [75]. Etiology is heterogeneous and comprises genetic, autoimmune, iatrogenic, and unknown causes. A careful evaluation of the medical history including medical treatments, previous surgery or hereditary disorders is recommended. Laboratory findings are an elevated FSH > 30 mIU/ml, determined twice with 4–6 weeks in between. AMH is of no benefit for the final diagnosis [1, 2].

All types of menstrual disturbances and various climacteric symptoms can be found. Women with POI/POF have a higher risk for estrogen-dependent diseases. However, there is no prospective study which shows lower mortality by hormonal substitution. It might make sense to continue hormonal treatment in women with POF until the average age of menopause.

There seems to be no difference between a hormonal substitution with HRT or contraceptive pills, although two studies reported a small difference in blood pressure in favour of HRT compared to contraceptive pills [76, 77]. Both treatment options can be discussed with POI/POF patients. In women > 40 years, HRT should be preferred.

Women with POI/POF should be informed about the following facts:

- HRT might have a positive effect on blood pressure compared to contraceptive pills.
- HRT and contraceptive pills have a positive effect on bone health.
- HRT does not offer protection from pregnancies.
- Women with POI and contraindications for HRT should be advised about cardiovascular risks, bone health and alternative treatments of climacteric symptoms.
- Women with POI should be generously transferred to specialized centres.
Informing women about HRT

Women should be informed about risks and benefits of medical interventions. The communication should include the probabilities of expected benefits and possible risks of harm with the patient and possibly with an accompanying person. For an individual assessment and evaluation of the probability of benefit and the risk of harm of HRT individual factors such as the woman’s general state of health, age at menopause, previous HRT, duration and use, doses and type of HRT and diseases while using HRT should be taken into consideration. To give adequate information about the risks to the woman seeking advice, the doctor must be familiar with the principles of risk calculation. He or she should also be able to communicate the probabilities in such a way that the patient can make her own individual decision for or against the initiation of HT. The figures necessary for this communication can be found in the long version and in Table 1 [1].

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Table 1 Estimated event rate difference associated with EPT or ET vs. placebo in postmenopausal women

| Events                  | EPT oral | ET oral |
|-------------------------|----------|---------|
| Breast cancer           | 9 (1 bis 19) | -7 (-14 bis 0.4) |
| Coronary heart disease  | 8 (0 bis 18) | -3 (-12 bis 8) |
| Stroke                  | 9 (2 bis 19) | 11 (2 bis 23) |
| Thromboembolism         | 21 (12 bis 33) | 11 (3 bis 22) |
| Dementia                | 22 (4 bis 53) | 12 (-4 bis 41) |
| Gallbladder disease     | 21 (10 bis 34) | 30 (16 bis 48) |
| Urinary incontinence    | 876 (606 bis 1168) | 1261 (880 bis 1689) |
| Bowel cancer            | -6 (-9 bis -1) | 2 (-3 bis 10) |
| Ovarian cancer          | 2 (-1 bis 6) | No data |
| Lung cancer             | 1 (-4 bis 7) | 1 (-4 bis 8) |
| Fractures (osteoporosis)| -44 (-71 bis -13) | -53 (-69 bis -39) |
| Diabetes                | -14 (-24 bis -3) | -19 (-34 bis -3) |
| Mortality               | 1 (-9 bis 12) | 1 (-10 bis 14) |

*a* If confidence interval includes 1, the result is not statistically significant

*b* Estrogen alone only in hysterectomised women

*c* For transdermal ET in doses from up to 50 µg/day observational studies did not demonstrate an influence of the risk for coronary heart disease, stroke or thromboembolism (modified table 3, Gartlehner et al. JAMA 2017)

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Compliance with ethical standards

Conflict of interest Author O. Ortmann declares that he has no conflict of interest. Author M. J. Beckermann declares that she has no conflict of interest. Author E. C. Inwald declares that she has no conflict of interest. Author T. Strowitzki declares that he has no conflict of interest. Author E. Windler declares that he has no conflict of interest. Author C. Tempfer declares that he has no conflict of interest.

Human/animal rights This article does not contain any studies with human participants or animals performed by any of the authors.

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Appendix

Detailed information on the development and methodology of the guideline following the criteria of the AWMF can be found on the AWMF homepage (see [1]).

Guideline group

Coordinating guideline author in charge.

| Author | Association of Scientific Medical Societies (AWMF) |
|--------|---------------------------------------------------|
| Prof. Dr. Olaf Ortmann | Deutsche Gesellschaft für Gynäkologie und Geburtshilfe (DGGG) |
|         | German Society of Gynecology                     |

Guideline group

DGGG-Working Group
AWMF/Non-AWMF-Scientific Society, Organisation

Arbeitgemeinschaft Gynäkologische Onkologie (AGO)
Arbeitgemeinschaft für Urogynäkologie und plastische Beckenbodenrekonstruktion (AGUB)
Berufsverband der Frauenärzte (BVF)
Statements and recommendations

Grade of recommendation:

A  strong recommendation.
B  recommendation with moderate obligation.
0  open recommendation with low obligation.

Degree of consensus:

+++  strong consensus (> 95% of delegates agree).
++  consensus (> 75 to 95% of delegates agree).
+   majority agrees (> 50 to 75% of delegates agree).
−   no consensus (< 51% of delegates agree).
LLA  Leitlinienadaptation, guideline adaptation.

*Dissenting opinion of the German Society for Phytotherapy (GPT):

The GPT does not support the undifferentiated recommendation on the use of cimicifuga, the level of evidence and grade of recommendation. In contrast to other
cimicifuga products (e.g., food supplements), cimicifuga medicinal products with marketing authorizations have proven their usefulness. Only these should be recommended.

Grade of recommendation:

A (isopropanolic Cimicifuga medicinal products).
B (ethanolic Cimicifuga medicinal products)

Level of evidence:

1b (isopropanolic Cimicifuga medicinal products).
2b (ethanolic Cimicifuga medicinal products)

For the full text on the position of the GPT see electronic supplementary material.

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