Femoston® 1/10. film-coated tablets

1 mg estradiol, and a combination of 1 mg estradiol and 10 mg dydrogesterone



Read this entire leaflet carefully before you start Keep this leaflet. You may need to read it again. If you

have questions not answered by this pamphlet, please ask your doctor or pharmacist. This medicine has been on to others. It may harm them, even if their symptoms

Femoston 1/10 is a round, film-coated tablet for ora administration Femoston 1/10 contains 14 white tablets, each of which

contains 1 mg estradiol and 14 grey tablets, each of which contains 1 mg estradiol and 10 mg dydrogesterone All tablets bear the inscription 379 on one side and S or

Excipients (non-medicinal ingredients):

mellose maize starch colloidal anhydrous silica. magnesium stearate

White tablet film-coating contains: titanium dioxide (E171) nypromellose, macrogol 400

Grey tablet film-coating contains: titanium dioxide (E171), iron oxide black (E172), polyvinyl alcohol, macrogol 3350,

emoston 1/10 is used as a Hormone Replacement Therapy (HRT) to treat symptoms of estrogen deficiency which are experienced by women in the years following menonause These symptoms vary from woman to woman and can include: hot flashes, night sweats, sleeping problems, vaginal dryness and urinary problems

Femoston 1/10 is also used to prevent bone thinning (osteoporosis) in post-menopausal women who are risk for bone fractures and who are intolerant of ontraindicated for other medicinal products approved for he prevention of osteoporosis.

Always take Femoston 1/10 exactly as your doctor has

prescribed. If you have any questions, you should ask you loctor or pharmacist. Do not start taking Femoston 1/10 until at least 12 months

after your last natural period. If you are currently not taking any HRT product or are

switching from a continuous combined preparation (i.e. both oestrogen and progesterone are taken daily in one tablet) you can start taking Femoston 1/10 on any menstrual spotting, start taking Femoston 1/10 on the first

If you are switching from a 'cyclic' or 'sequential' HRT product (this is when you take an oestrogen tablet or use a patch for the first part of your cycle, followed by a daily ablet containing both an oestrogen and a progestor up to 14 days) start taking Femoston 1/10 the after you finish the previous pack (i.e.: at the end of the progestogen phase).

If you are changing from a previous seguential hormone replacement therapy, your menopausal status may not be known. Also, in some women endogenous gestroger may still be produced. This could result in unpredictab bleeding patterns, i.e. you may experience breakthrough

The sequence in which to take your tablets is clearly indicated on the blister Specifically take one white tablet daily for the first 14 days of a 28 day cycle and one Always take Femoston 1/10 continuously without a break

Femoston 1/10 can be taken with or without food; however the tablet should be swallowed with water.

Try to take your tablet at the same time each day. This will ensure that there is a constant amount of the product n your body. This will also help you to remember to take vour tablets

If you have forgotten to take a tablet it should be taken as soon as possible. If more than 12 hours have elapsed, you should take the next tablet without taking the forgotten one Do not take a double dose. Be advised that breakthrough

Regardless of whether you are starting or continuing therapy for postmenopausal symptoms, your doctor will always prescribe the lowest possible dose for the shortest period of time (see section "Warnings and special precautions for use").

general, your doctor will start your treatment with on 1/10. Your dosage may be adjusted thereafter depending on your response to the therapy. If your (postyour doctor may increase the dosage by prescribing you If you are taking Femoston to prevent osteoporosis, your

doctor will adjust the dose individually according to your hone mass

Do not stop taking Femoston without first talking to your

The experience in treating women older than 65 is limited. Femoston 1/10 is not indicated for the use in children.

Contraindications Do not take Femoston 1/10 if:

■ you are allergic (hypersensitive) to estradiol, dydrogesterone to any of the other ingredients of Femoston (see

you have, have had or your doctor suspects you may you have or your doctor suspects you may have a tumour

poestrogen-dependent (such as cancer of the uterine lining (endometrial cancer))
oor that is progestogen-dependent (such as meningioma)

■ you have undiagnosed genital bleeding (i.e. unclear vou have abnormal thickening of the lining of the uterus endometrial hyperplasia) for which you have not yet

wou have or have had a blood clot(s) in your leg(s) or lungs, for which no obvious cause has been found venous thromboembolism i.e.; deep venous thrombosis

■ vou have or recently have had a disease caused by blood clots in the arteries (arterial thromboembolic disease) such as angina or a heart attack (myocardial infarction

function test values have not vet returned to normal you have a rare blood pigment disorder called "porphyria which may be either passed down in families (inherited) or acquired.

Warnings and special precautions for use

For the treatment of postmenopausal symptoms, treatment with Femoston 1/10 should only be started if your symptoms seriously affect your quality of life. In all cases your doctor will carefully consider both the risks and should only be continued as long as the benefits outweigh the risks. Annual re-evaluations are recommended.

Medical examination and follow-up
Refore you start or restart Hormone Replacement Therapy (HRT), your doctor will ask you for a complete personal and family medical history. According to the findings, your doctor will perform a full examination, possibly including a pelvic and breast examination. Your doctor will also take nto account any contraindications and warnings for use

including regular breast screenings (mammography) according to your doctor's recommendations and depending on your personal situation, but at least once a year

Important note: Do regular self-breast examinations. If vou notice any changes in your breasts tell your doctor mediately. If you are not sure how to do a self-breast examination or what changes to look for, ask your doctor.

Conditions which need supervision Your doctor will closely supervise you if you have or have had any of the following conditions, or if pregnancy or evious hormone treatment has worsened the condition. It s possible for these conditions to recur or to be aggravated n treatment with Femoston 1/10 in particula

aberrant growth of the lining of the uterus (uterine fibroids (leigmygma)) or of uterine tissues outside the uterus (endometriosis) ary of arrisk factors for blood clots ar other disorder caused by the blockage of blood vessels (thrombo embolic disorders) (see "Venous thromboembolism

an increased risk for nestrogen-dependent tumours e.g. a direct (1st degree, such as a mother or a sister) elative with breast cancer

high blood pressure (hypertension) liver disorders, e.g. adenoma, which is a benign tumour diabetes mellitus, with or without concurrent vascular

gall stones (cholelithiasis Migraina or cavara haadacha

an immune system disorder affecting many organs of the body (systemic lupus erythematosus)

a history of abnormal thickening of the uterine lining

ndométrial hyperplasia) (see bělow) seizures (epilépsy)

inner ear disease (otosclerosis) Reasons to stop taking Femoston immediately; Your doctor will stop your therapy with Femoston 1/10 if

any of the contraindications apply to you or if he notices ■ vellowing of the skin and/or whites of your eyes (iaundice)

= significant increase in your blood pressure ■ new onset of migraine-type headache = pregnancy

Important note: If you notice any of the above listed conditions stop taking Femoston immediately and talk to

ping abnormal overgrowth of the tissues lining the uterus (endometrial hyperplasia) and cancer (carcinoma) is increased when oestrogens are administered alone for prolonged periods (see section "Undesirable effects"). The addition of a progestogen (separately or in a combined tablet) for at least 12 days per cycle greatly reduces this risk in women with an intact uterus (non

nexpected bleeding (breakthrough bleeding) and spotting may occasionally occur during the first months of treatment. If you experience breakthrough bleeding or snotting after you have been on the therapy for some stopped, inform your doctor immediately. Your doctor will vestigate the cause of the bleeding and may perform tests (e.g. a uterine (endometrial) biopsy) to rule out iterine cancer (endometrial malignancies).

ireast cancer possible link between treatment of women with hormones and the development of breast cancer. Results are as

A randomicad placeho-controlled trial the Women's Health Initiative study (WHI), and epidemiological studies, including the Million Women Study (MWS), have reported an increased risk of breast cancer in women taking oestrogens nestrogen-progestogen combinations or tibolone for HRT for several years (see section "Undesirable effects"). For all types of HRT, there is an increased risk of developing breast cancer after several years of continuous use. This extra risk increases the longer HRT is continued, but eturns to normal levels within a few (at most five) years after stopping treatment.

The MWS reported an increased risk of developing breas cancer when a combined (oestrogen-progestogen) product was used regardless of the type of progestogen used, the method of administration (sequential or continuous) or the route of administration

In the WHI study the continuous combined conjugated equine oestrogen and medroxyprogesterone acetate (CEE+MPA)) product used was associated with breast cancers that were slightly larger in size and more frequently had local lymph node metastases compared to women not taking HRT.

especially oestrogen-progestogen combined products, increases the density of mammographic images which may make the detection of breast cancer

Venous thromboembolism

HRT may increase your risk of developing blood clots in the veins of the legs or lungs (venous thromboembolism

studies found the risk to be two to three times higher women taking HRT compared to women not taking HR non-users it is estimated that the number of cases of VTF that will occur over a 5 year period is about 3 per 1006 women aged 50-59 years; and 8 per 1000 women aged between 60-69 years. It is estimated that in healthy women who have used HRT for at least 5 years, the number of additional cases of VTF over a 5 year period will be een 2 and 6 (best estimate= 4) per 1000 women age 59 years; and between 5 and 15 (best estimate= 9) pe 1000 women aged 60-69 years. The probability of such a romboemblism occurring is higher during the first year of HRT as opposed to later

In general the risk of developing VTE while on Femostor 1/10 is increased if you are severely obese (Body Mass Index > 30 kg/m²) or if you have the immune system disorder "systemic lunus erythematosus" It is unclear whether varicose veins contribute to the risk of VTE. If you have varicose veins please inform your

doctor before starting HRT.

You may be predisposed to develop VTE if you, or a family member, have or have had a history of VTE or other known disease which causes blood clots. HRT may increase this risk. Your doctor will investigate any personal or strong family history of blood clot disorders (thromboembolism) or recurrent miscarriages in order to be sure you are not predisposed to VTE or, alternatively, to assess the potential risks of VTE to you. You will not be started on Femoston 1/10 until a thorough evaluation of these factors has been made or you have started taking blood thinning medicines (anticoagulants). Also, if v are already taking an anticoagulant, talk to your doctor

You will only be prescribed Femoston 1/10 if the benefits of HRT far outweigh the risks of developing VTE. The risk of VTE may be temporarily increased if you have been immobile (e.g. bed ridden or in a wheelchair) for a prolonged period, if you have suffered a major trauma or have had a major surgery. As is always done following surgery, your doctor will do everything possible to heli prevent a VTE from occurring. If you are scheduled for an elective surgery which is likely to result in prolonged immoilization, such as belly (abdominal) or leg (orthopaedi surgery, your doctor may temporarily stop your HRT four to six weeks before the procedure. The treatment should not be restarted until you are fully recovered from

f you develop a VTE after starting therapy with Femostor 1/10, your doctor will stop your therapy. Furthermore, stop taking Femoston 1/10 and contact your doctor mmediately if you develop any potentially thror symptoms such as: painful leg swelling, sudden ches ain, and/or difficulty breathing (dyspnea).

Coronary artery disease (CAD) no benefits with the use of a specific type of HRT continuous combined conjugated oestrogens and nedroxy progesterone acetate (MPA)) to the cardiovascular el) system. Two large clinical trials (WHI and HERS, i.e. Heart and Oestrogen/progestin Replacement Study) showed that the risk of suffering (morbidity) from a cardiovascular disease may be higher in the first year of and that there are no overall cardiovascular be here is not enough information concerning other types of HRT to determine if these findings also extend to other HRT products (including Femoston 1/10)

rding to the Women's Health Initiative trial (WHI-trial) the risk to healthy women for ischaemic stroke (resulting from a deficiency of blood supplied to the brain) is higher when taking HRT which contains continuous combined conjugated oestrogens and MPA.

As comparison: for those women not treated with hormone occur over a 5 year period is about 3 per 1000 women years. In women who have been treated with conjugated oestrogens and MPAs for 5 years, the number of addition cases of stroke will be between 0 and 3 (best estimate=

per 1000 users aged 50-59 years and between 1 and 9 (best estimate= 4) per 1000 users aged 60-69 years. It is unknown whether the increased risk also extends to other HRT products (including Femoston 1/10).

term (at least 5-10 years) use of nestrogen-only HR Long-term (at least 5-10 years) use or desired. The products in women whose uterus has been removed a specified with an increase. of ovarian cancer in some epidemiological studie It is uncertain whether long-term use of combined HRT (such as Femoston 1/10) confers a different risk than oestrogen-only products.

ogens may cause fluid retention, so your doctor will monitor you carefully if you have any type of heart or kidney disease. Furthermore, if you have seshould be monitored closely by your doctor, since thi condition can cause an increase of circulating active ingredients of Femoston 1/10 in your blood.

you have a high concentration of lipids in your blood (hypertriglyceridemia), you should visit your doctor more frequently while on HRT (whether you take an oestogenonly or combined product). In rare cases large increases of blood lipid levels (triglycerides) leading to inflammation of the nancreas have been reported with pestronen herapy in patients with this condition.

Destrogens may affect thyroid gland function. Talk to your

doctor if you have a thyroid gland disease or problen before you start taking HRT.

Information for the doctor: Specifically, oestrogens increase thyroid binding globulin (TBG), leading to increased circulating total thyroid hormone, as measured by protein-bound doline (PBI), 74 levels (by column or by radio-immunoassay) or T3 levels (by radio-immunoassay) T3 resin untake is decreased reflecting the elevated TBG. Free T4 and free T3 concentrations are unaltered. Other binding proteins may be elevate in serum, i.e. corticoid binding globulin (CBG), sex-hormone binding globulin (SHBG) leading to increased culating corticosteroids and sex steroids, respectively ree or biologically active hormone concentrations are unchanged. Other plasma proteins may be increase (angiotensinogen/renin substrate alpha-l-antitrypsin

There is no conclusive evidence that HRT improves the ability to think clearly (cognitive function). There is some evidence from the WHI trial of an increased risk of de mentia in women who start using a specific type of HRT (continuous combined CEE and MPA) after the age of 65. It is unknown whether the findings apply to younger post-menopausal women or other HRT products (including

Do not take this medicine if you have any of the following are hereditary problems: galactose intolerance, the Lapp lactase deficiency or glucose-galactose malabsorption Femorton 1/10 is not a contracentive and is not intended to be used by women who could become pregnant. In case

of doubt, use a non-hormonal contraceptive Interactions with other medications Please tell your doctor or pharmacist if you are taking r have recently taken any other medicines including

medicines obtained without a prescription and herba Please make sure to read the leaflet of any other medicine you are taking at the same time as Femoston.

No studies have been performed to investigate interactions between Femoston 1/10 and other medicina

The following may reduce the effects of Femoston and give rise to bleeding or spotting: Medicines for the treatment of: epilepsy (such as phenobarbital, carbamazepine and

oHIV infection [AIDS] (such as ritonavir, nelfina m herbal remedies containing St John's wort (the extract of the plant called St. John's wort is included in certain herbal preparations used particularly for menopausa

Destrogens might slow the breakdown of other drugs which may lead to dangerously high levels of such drugs in the blood. Therefore, careful drug monitoring and possibly a dosage decrease may be necessary cyclosporin A. and theophylline.

Information for the doctor

e efficacy of oestrogens and progestogens might be

 The metabolism of oestrogens and progestogens may be increased by concomitant use of substances known to induce drug-metabolising enzymes, specifically the P450 enzymes 2B6, 3A4, 3A5, 3A7, such as anticonvulsants phenobarbital, carbamazepine and phenytoin and anti-infectives (e.g. rifampicin, rifabutin, nevirapine

inhibitors of CYP450 3A4, A5, A7, by contrast exhibit nducing properties when used concomitantly with steroid Gastrointestinal disorder Common: Nausea, abdominal pain, flatulence ■ Herbal preparations containing St. John's Wort (Hypericum

perforatum) may induce the metabolism of oestrogens and progestogens via the CYP450 3A4 pathway. Clinically, an increased metabolism of oestrogens and progestogens may lead to decreased effect and changes in the uterine bleeding profile.

Destrogens might interfere with the metabolism of other ens per se may inhibit CYP450 drug-metabolising

enzymes via competitive inhibition. This is in particular to e considered for substrates with a narrow therapeutic index such as acrolimus and cyclosporine A (CYP450 3A4, 3A3)

theophylline (CYP450 1A2). Clinically this may lead to a plasma increase of the affected ring for an extended period of time might be necessary and

and theophylline may be necessary. Pregnancy and lactation

Ask your doctor or pharmacist for advice before taking any medicine during pregnancy

Important: Do not take Femoston 1/10 if you are pregnant Femoston 1/10 is for use in post-menopausal women onl

If you become (or think you are) pregnant while being treated with Femoston 1/10, stop taking the medicine immediately and tell your doctor. The results of most epidemiological studies concerning the accidental exposure of a foetus to oestrogen/prod

combinations show no negative effects to the developing baby (no teratogenic or foetotoxic effects). Effects on shility to drive and use machines emoston 1/10 has no or negligible influence on the ability to drive and use machines.

Important information about the ingredients iston 1/10 contains lactose monohydrate. If you have peen told by your doctor that you have an intolerance o some sugars, especially lactose, contact your docto efore taking this medicinal product.

Undesirable effects Like all medicines, Femoston 1/10 can cause side effects, although not everybody experiences them. If you notice any side effects not mentioned in this leaflet

or if any of the side effects gets serious, please inform your doctor or pharmacist. Undesirable effects reported in clinical trials and in postmarketing experience are the following:

The frequencies of study related side effects are ranked according to the following: common (frequency 1-10%) uncommon (frequency 0.1-1%), rare (frequency 0.1-1%), very rare (frequency <0.01%, including isolated

Undesirable Effects by System Organ Class: Uncommon: Vaginal yeast infections (vaginal candidiasis) Neoplasms benign, malignant and unspecified

Uncommon: Increase in size uterine fibroids (leiomyoma Blood and lymphatic system disorders Very rare: Illness due to the destruction of red blood cells (haemolytic anaemia) symptoms may include paleness of the skin, generalized weakness and/or difficulty breathing

Immune system disorders Very rare: Allergic reactions (hypersensitivity) Psychiatric disorders

Uncommon: Depression, changes in sex drive, nervousness Nervous system disorders Common: Migraine, headache Uncommon: Dizziness

Very rare: Involuntary muscle twitches (chorea)

n/bending of the membrane covering the eve (steepening of corneal curvature), intolerance to contact

Cardiac disorders Very rare: Heart attack (myocardial infarction)

Vascular disorders Uncommon: Blood clots in the legs or lungs (venous thromboembolism (see below for further information)) Very rare: Stroke

Very rare: Vomiting Henatohiliary disorders

non: Gall bladder disease are: Abnormal liver function, occasionally with vellowing of the skin, gums and/or inner eye membrane (jaundice weakness (asthenia) or general malaise, and abdominal

Skin and subcutaneous tissue disorders mon; Allergic skin reactions (e.g. rash, hives (urticaria),

tching (pruritus)) Very rare: Swelling of the limbs, face or throat, which may cause difficulty breathing (angioedema), red or brown patches on the skin (erythema multiforme/nodosum purplish patches or spots on the skin (vascular purpura kin discolouration, which may persist when drug is discontinued (chloasma or melasma)

Musculoskeletal and connective tissue disorders Common: Leg cramps Uncommon: Back pain

Reproductive system and breast disorders Common: Breast pain/tenderness non-menstrual uterine eding or spotting (metrorrhagia) and post-menopausal spotting, pelvic pair

non: Erosion of the lining of the cervix (uterine cervical erosion), cervical discharge, painful menstruation Rare: Breast enlargement, pre-menstrual syndrome

Very rare: Worsening of a rare blood pigment disorder called "porphyria" General disorders and administration site reactions Common: Muscle weakness (asthenia)

Uncommon: Swelling of the limbs (peripheral oedema) Common: Increase or decrease in weight

Congenital and familial/genetic disorders

reast cancer ased on results of a large number of epidemiological studies and one randomised placebo-controlled trial, the Women's Health Initiative (WHI), the overall risk of breast cancer increases the longer a woman is on HRT. This is true, both for women currently using HRT and those having

used HRT recently. *Below you will find detailed information concerning the risks associated with using HRT and developing breast cancer. For clarifications or further information, please consult your doctor.

For oestrogen-only HRT, estimates of relative risk (RR) from a reanalysis of original data from 51 epidemiological studies (in which more than 80% of HRT use was oestrogen-only HRT) and from the epidemiological Millio Women Study (MWS) are similar at 1.35 (95% CI: 1.21-1.49) and 1.30 (95% CI: 1.21– .40), respectively

For oestrogen-progestogen combined HRT, several epidemiological studies have reported an overall higher risk for breast cancer than with oestrogens alone.

The MWS reported that compared to never-users the use The MWS reported that, compared to never-users, the use of various types of oestrogen-progestogen combined HRT was associated with a higher risk of breast cancer (RR = 2.00, 95 % Cl: 1.88–2.12) than use of oestrogens alone (RR = 1.30, 95% Cl: 1.21–1.40) or use of libolone (RR = 1.45: 95% CI: 1.25-1.68).

The WHI trial reported a risk estimate of 1.24 (95% CI: 1.01-1.54) after 5.6 years of use of oestrogen-progestogen combined HRT (CEE + MPA) in all users compared with

The absolute risks calculated from the MWS and the WHI trials are presented below: The MWS has estimated, from the known average incidence of breast cancer in developed countries, that ■ for women not using HRT, about 32 in every 1000 are

expected to have breast cancer diagnosed between the ages of 50 and 64 years.

In the for 1000 current or recent users of HRT, the number of

additional cases during the corresponding period will be ofor users of oestrogen-only replacement therapy between 0 and 3 (best estimate = 1.5) for 5 years use between 3 and 7 (best estimate = 1.5) for 10 years use.

ofor users of oestrogen-progestogen combined HRT. between 5 and 7 (best estimate = 6) for 5 years' use between 18 and 20 (best estimate = 19) for 10 years' The WHI trial estimated that after 5.6 years of follow-up

of women between the ages of 50 and 79 years, ar additional 8 cases of invasive breast cancer would be due to oestrogen-progestogen combined HRT (CEE + MPA) per 10,000 women years. ording to calculations from the trial data, it is estimated

For 1000 women in the placebo group, about 16 cases of

invasive breast cancer would be diagnosed in 5 years. combined HRT (CFF + MPA), the number of additional 5 years' use. The number of additional cases of breast cancer in women

who use HRT is broadly similar for women who start HRT 5 - 65) (see section "Warnings and special precautions

Endometrial cancer

In women with an intact uterus, the risk of developing an abnormal growth of the lining of the uterus (endometrial ia) and cancer of the inner lining of the uterus (endometrial cancer) increases the longer a woman takes unopposed oestrogens. Results of epidemiological studies show that, for women not using HRT, approximately n every 1000 are expected to have endometrial cancer diagnosed between the ages of 50 and 65. Depending or the duration of treatment and oestrogen dose, the reporte increased risk for endometrial cancer among unoppose oestrogen users varies from 2- to 12-fold greater compared with non-users. Adding a progestogen to oestrogen-only therapy greatly reduces this increased risk. Venous thromboembolism, i.e. blood clots in the legs

pelvic or lungs (deep leg or pelvic venous thrombosis and pulmonary embolism), is more frequent among HRT users than among non-users. For further information, see sections "Contraindications" and "Warnings and special precautions for use".

Other adverse reactions that have been reported in ciation with oestrogen-progestogen treatment: ■ Tumors/neoplasms, benign, malignant and unspecified: oOestrogen-dependent neoplasms both self-limiting (benign) and invasive (malignant), e.g. endometrial cancer

olncrease in size of progestogen-dependent neoplasms (e.g. meningioma)

Immune system disorders: Systemic lupus erythematosus ■ Nervous system disorders: Possibility of developing

dementia, worsening of epileptic symptoms Vascular disorders: blood clots in the arteries (arterial

■ Benal and urinary disorders: loss of bladder control (urinary incontinence)

No case of overdose has been reported for Femoston 1/10 Both estradiol and dydronesterone are substances with unlikely to do any harm. However, symptoms of overdose nay include: nausea, vomiting, sleepiness and dizzines It is unlikely that any treatment will be necessary, however you (or someone else) take too many tablets inform you doctor immediately.

The above information is also applicable to cases of overdose in children **Pharmacodynamics** harmacotherapeutic group: Genito urinary sytem and

sex hormones, progestogens and oestrogens, sequential • The following is a detailed description of how the active ingredients (estradiol and dydrogesterone) of Femoston 1/10 work. For clarifications or further information please

The active ingredient, estradiol, is chemically and biologically identical to the endogenous human estradiol

and is, therefore classified as a human oestrogen, Estradiol the primary oestrogen and the most active of the ovarian normones. Endogenous oestrogens are involved in certain functions of the uterus and accessory organs, including the proliferation of the endometrium and the cyclic changes n the cervix and vagina.

Oestrogens are known to play an important role on bone and fat metabolism. Furthermore, oestrogens also affect the autonomic nervous system and may have indirect positive psychotropic actions

Dydrogesterone is an orally-active progestogen having an activity comparable to parenterally administered rogesterone n the context of HRT, dydrogesterone produces a complete secretory endometrium in an oestrogen-primed uterus hereby providing protection for oestroge

increased risk for endometrial hyperplasia and/or carcinogenesis, without androgenic side-effects oestrogens promote the growth of the endometrium inopposed oestrogens increase the risk of endometria hyperplasia and cancer. The addition of a progestoger

rplasia in non-hysterectomised women. · Relow find information on the results of clinical trials for Femoston products for clarifications or further information please consult your doctor. Clinical trial Information

f oestrogen-deficiency symptoms and bleeding

Belief of menopausal symptoms was achieved during the first few weeks of treatment. Regular withdrawal bleeding with Femoston 2/10 occurred in approximately 909 of women with a mean duration of 5 days. Withdrawal bleeding usually started on the day of the last tablet of the progestogen phase. Breakthrough bleeding and/or spotting appeared in approximately 10% of the women. orrhoea (no bleeding or spotting) occurred in 15% of the women per cycle during the first year of With Femoston 1/10, 75 - 80% of women had regular

withdrawal bleeding. The start day and duration of blee and the number of women with intermittent bleeding was the same as with Femoston 2/10, but there were more women without any bleeding per cycle (10 - 25% per cycle).

Prevention of osteoporosis: Destrogen deficiency at menopause is associated with

sing bone turnover and decline in bone mass. The effect of oestrogens on the bone mineral density is dose-dependent. Protection appears to be effective for as long as treatment is continued. After discontinuation of HRT, bone mass is lost at a rate similar to that in

untreated women.

Evidence from the WHI trial and meta-analysed trials ws that current use of HRT, alone or in combinati with a progestogen - given to predominantly healthy women - reduces the risk of hip, vertebral, and other osteoporotic fractures. HRT may also prevent fractures in women with low bone density and/or established osteoporosis, but the evidence for that is limited

After two years of treatment with Femoston 2 increase in lumbar spine bone mineral density (BMD was 6.7% ± 3.9 % (mean ± SD). For Fernoston 1/10 the percentage of women who maintained or gained BMD in lumbar zone during treatment was 94.5%. For Femoston 1/10 the increase in lumbar spine BMD was with no change or an increase in lumbar spine BMD was 93.0%. Femoston also had an effect on hip BMD. The increase after two years of treatment with 1 mg estradiol was 2.7% ± 4.2 % (mean ± SD) at femoral neck, 3.5% ± 5.0% (mean + SD) at trochanter and 2.7%+6.7% (mean SD) at Wards triangle, after two years of treatment with mg estradiol these figures where respectively 2.6% ± %; 4.6% ± 5.0% and 4.1% ± 7.4% women who maintained or gained BMD in the 3 hip areas after treatment with 1 mg estradiol was 67-78% and 71-88% after treatment with 2 mg estradiol.

Pharmacokinetics

The following is a detailed description of how the active ingredients (estradiol and dydrogesterone) of Fernoston 1/10 are processed by your body. For clarifications or further information please consult your doctor. oral administration, micronized estradiol is readily

absorbed, but extensively metabolised. The major

unconjugated and conjugated metabolites are estrone

and estrone sulphate respectively. These metabolites car contribute to the oestrogen activity, either directly or after conversion to estradiol. Estrone sulphate may undergo enterohepatic circulation. In urine, the major compounds are the ducuronides of estrone and estradiol Oestrogens are secreted in the milk of nursing mothers

<u>Jydrogesterone</u>
After oral administration of labelled dydrogesterone, on verage 63% of the dose is excreted into the urine. Within 72 hours excretion is complete. In man, dvdrogesterone is

The main metabolite of dydrogesterone is 20 α-dihydro redominantly as the glucuronic acid conjugate. A common eature of all metabolites characterized is the retention of the 4,6 diene-3-one configuration of the parent compound and the absence of 17α -hydroxylation. This explains the lack of oestrogenic and androgenic effects of dydro

gesterone.

After oral administration of dydrogesterone, plasma concentrations of DHD are substantially higher as compared to the parent drug. The AUC and Cmax ratios f DHD to dydrogesterone are in the order of 40 and 25

production based on pregnanediol excretion therefore

Dydrogesterone is rapidly absorbed. The T_{max} values of dydrogesterone and DHD vary between 0.5 and 2.5 hours. Mean terminal half lives of dydrogesterone and DHD vary between 5 to 7 and 14 to 17 hours, respectively. Dydrogesterone is not excreted in urine as pregnanediol gesterone. Analysis of endogenous progesterone

remains possible. After oral administration, the T_{max} values for dydrogesterone and DHD were 1.25 hours and 1.21 hours respective The mean terminal half times of dydrogesterone and DHI were was 6.56 hours and 17.62 hours respectively.

Incompatibilities Not applicable

Shelf life and storage conditions Do not store above 30°C

Manufactured b

The Netherlands

the medication

Store in the original nackage Do not use this medicine after the expiry date stated on

Keep this medicine out of the reach and sight of children

Pack sizes
28. 84 or 280 (10 x 28) film-coated tablets per pack (not all ack sizes may be marketed) The blisters are made of PVC/PVDC or PVC with a

covering of aluminium foil Further information Any unused product or waste material should be disposed of in accordance with local requirements.

The information in this leaflet is limited. For further nformation, please contact your doctor or pharmacist Date of information

bbott Healthcare Products B.V..

THIS MEDICATION is a product which affects your health and its use contrary to instructions is dangerous to you. Strictly follow the doctor's prescription, the method of use and the instructions of the pharmacist who sold you

The doctor and the pharmacist are the experts medicines, their henefits and risks Do not interrupt the period of treatment prescriber

Do not repeat the same prescription without first consulting your doctor. Keep all medications out of reach of children

Council of Arab Health Ministers Union of Arab Pharmacists.

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