

The Effect of Hormone Replacement Therapy in Menopause on Cardioprotection

POSTMENOPOZAL HORMON REPLASMAN TEDAVİSİNİN
KARDİYOPROTEKTİF ETKİLERİ

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SUMMARY

Cardiovascular disease (CVD) is the leading cause of death women, and after menopause the incidence increases rapidly. The premenopausal state and estrogen status, appear to be a prophylactic against the mortality risk from CVD. The protective effect is believed to be mediated by beneficial changes in cholesterol levels. Estrogen decreases low density lipoprotein cholesterol (LDL-C) and increases high density lipoprotein cholesterol (HDL-C). The other possible mechanism is the direct effect of estrogen on arterial intima. The type and route of estrogen used in hormone replacement therapy determines the positive and negative effect of estrogen on the cardiovascular system. Most studies show a 50% or greater reduction in CVD and related mortality with postmenopausal estrogen administration. Progestogen addition to hormone replacement, may attenuate the beneficial effects of estrogen on cholesterol. However if used in low doses progesteron may not exert this negative effect on the cardiovascular system.

Key Words: Estrogen, Cardiovascular disease

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Women rarely have heart disease until after menopause unless they have a major coexisting disease such as familiar hypercholesterolemia or diabetes. After menopause the Incidence of cardiovascular disease accelerates rapidly. In the Framingham study iur example, the combined fatal and non-fatal rates of cardiovascular disease in women aged 45 to 49 lag behind those in men by about 15 years. This lag de-

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ÖZET

Kardiovasküler hastalıklar kadınlarda en önde gelen ölüm nedenidir ve menopoz sonrası insidansta hızlı bir artış gözlenmektedir ve östrojenin kardiovasküler hastalıklar açısından profilaksi oluşturduğu düşünülmektedir. Koruyucu etkinin kan kolesterol düzeylerinde oluşan olumlu değişiklikler sonucu ortaya çıktığına inanılmaktadır. Östrojen düşük dansiteli lipoproteinleri azaltırken yüksek dansiteli lipoprotein düzeylerinin artmasına neden olur. Diğer bir muhtemel mekanizma da östrojen hormonunun arteryal intima üzerindeki direkt etkisidir. Hormon replasman tedavisinde kullanılan östrojenin tipi ve uygulama yolu kardiovasküler sistem üzerindeki olumlu ve olumsuz etkileri belirler. Pekçok çalışmayla postmenopozal östrojen uygulamasının kardiovasküler hastalık ve buna bağlı mortalityde %50 ya da daha fazla azalma sağladığı gösterilmiştir. Östrojen replasman tedavisine progesteron eklenmesini östrojenin kolesterol üzerindeki faydalı etkilerini zayıflatabilir. Yine de düşük dozlarda kullanılan progesteron kardiovasküler sistem üzerinde olumsuz etkiler göstermeyebilir.

Anahtar Kelimeler: Östrojen, Kardiovasküler hastalık

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creases to 10 years at ages 70 to 75 and after 75 the rates are nearly equal (1). In most industrialized countries, women can expect to live more than one third of their lives in postmenopausal state. Cardiovascular disease is the leading cause of death in women and accounts for 52% of all deaths in U.S. women (2). Coronary artery disease alone effects more than two million women per year in the U.S.A. Within the first several years after natural menopause, women have no appreciably increased risk of cardiovascular disease (3). However, women who have premature menopause or oophorectomy are at substantially increased risk of cardiovascular disease. Some studies have suggested that the risk in these women is seven times increased

than in women with intact ovaries (4). The risk is higher in women with premature menopause because estrogen deprivation appears earlier and more abruptly.

The premenopausal state and estrogen status, appear to be a prophylactic against the mortality risk from cardiovascular disease. The purpose of this article is to review the literature to determine the mechanism of hormonally related cardioprotection.

MECHANISMS OF ESTROGEN RELATED CARDIOPROTECTION

The Framingham data (1) revealed no substantial differences between men and women in known risk factors such as elevated blood pressure, impaired glucose tolerance, except increased total cholesterol with advancing age. Before menopause, women have lower cholesterol levels than men, but their levels start to increase with onset of menopause, and increase to a greater extent than do cholesterol levels in aging men (5). Epidemiologic data from the Lipid Research Clinics (6) have revealed that for every 1% increase in total cholesterol results in 2% increase in the risk of myocardial infarction. The pathogenetic role of cholesterol in is coronary heart disease therefore significant.

There is still dispute as to whether the principal injury to the arterial intima that culminates in myocardial infarction is primarily of a lipid or nonlipid origin. It has been established that all cases of impeding myocardial infarction involve platelet aggregation and release of growth promoting substances such as platelet derived growth factor and transforming growth factor beta (7). These chemotactic factors entrapping lipids can induce the formation of foam cells. Foam cells are lipid carrying and processing macrophages that have been activated to engulf lipid, and specifically, LDL cholesterol. Damage results because the foam cells eventually occupy the lumen of the coronary vessel and weaken the vessel wall. Rupture of arterial wall will activate the clotting mechanism and development of a coronary thrombus will lead to myocardial infarction. Macrophages may further contribute to atherogenesis by oxidation of LDL cholesterol (8,9). The oxidant forms appear to have different receptor attachments and can not be appropriately removed, resulting in accumulation of the cholesterol in the vessels. These features make the oxidant forms of LDL cholesterol extremely atherogenic. The LDL receptor is therefore critical for removal of cholesterol from the circulation (10). If the LDL receptors are abnormal or diminished in number, severe atherosclerosis will result. Deficiency of LDL receptor can be absolute, from genetic causes or relative, in proportion to excessive dietary cholesterol intake. Levels of LDL cholesterol significantly affect cardiovascular disease mortality; for every 11% reduction in LDL cholesterol levels there is a 19% reduction in coronary artery disease risk (11).

High density lipoprotein (HDL) cholesterol plays major role in the collection of excessive cholesterol and its transport back to liver for degradation and reformation other lipoproteins. There are several forms of HDL cholesterol, but the one that is most active in reverse cholesterol transport is HDL 2 (12). The HDL2 particles are catabolized primarily in the liver under the influence of hepatic lipase. Hepatic lipase activity decreases during estrogen treatment and increases with progesterone treatment. While estrogen shuts down hepatic lipase activity, HDL cholesterol can increase in the circulation and continue to facilitate increased cholesterol transport. In both men and women, extrapolation of these data shows a greater number of coronary vessels affected when the HDL cholesterol level is lower (13). Interestingly, increased cardiovascular risk after menopause is not a result of low HDL cholesterol levels. Cross sectional data have failed to show a major change in HDL cholesterol as a function of age during menopause. However HDL cholesterol levels do decline in premenopausal women made acutely estrogen deficient or in women undergoing oophorectomy. Therefore a decline in HDL cholesterol per se do not seem to increase cardiovascular risk in menopause. The causal factor changing during menopause is total cholesterol, 70% of which is composed of LDL cholesterol. Although HDL cholesterol levels does not tend to decline by postmenopausal period, it is supposed that estrogen replacement can cause an increase in HDL cholesterol in a postmenopausal woman. It was reported that 1.6 mg/dL increase in HDL cholesterol would predict a 9.2% reduction in coronary artery disease. The actual observed reduction was 7% (14).

Estrogen seems to increase local production of prostacyclin which has vasodilatory properties and anti-aggregant effects. It is well known that estrogen induces an increase in blood flow. A favorable thromboxan / prostacyclin ratio plus increased blood flow may help explain the cardioprotective effects of estrogen. It is estimated that as much as 50% of the cardioprotection by estrogen in postmenopausal women can not be explained by estrogen induced increase in HDL cholesterol levels. This suggestion directed the investigators to assess local action of estrogen on vessels. Estrogen and progesterone receptors have been found in arterial endothelial and smooth muscle cells of several mammalian species (15). Other studies have shown that estrogen treatment in vivo or in vitro is associated with reductions in lipoprotein - induced arterial smooth muscle cell proliferation (16), inhibition of the myointimal proliferation associated with mechanical endothelial injury (17), decreased collagen and elastin production (18), increased collagen and elastin degradation (19) and increased prostacyclin production (20) by arterial smooth muscle cells. These studies indicate that vascular estrogen receptors have functional capabilities and may determine atherosclerotic response at the level of arterial intima.

ESTROGEN REPLACEMENT AND CARDIOVASCULAR DISEASE

More than 20 studies have compared the frequency of myocardial infarction or ischemic heart disease according to estrogen use (Table 1). Only two (29,40) of these studies suggested increase of risk with hormone use in postmenopausal women and two others showed no risk or benefit (28,41). All other studies suggest significant protection with a risk reduction of at least 50% in most cases. The Framingham study found an elevated risk which was not statistically significant when women with angina were omitted (40). As a consequence this study involved a small number of patients and had imprecise disease end points. A subsequent reanalysis of this data showed a nonsignificant protective effect among young women but a nonsignificant adverse effect among older women (41). There is less benefit perhaps an adverse effect among women taking more than 1.25mg of conjugated estrogen daily. Such high doses were common in Framingham study which may partly explain their discrepant results.

In the Lipid Research Clinic study (35) the relative risk of cardiovascular disease was lower in hormone using women in each age group compared with nonusing women. Henderson et al (38) also found a 50% lower risk of heart disease in women older than 70 who were treated with estrogen, suggesting that the protection continuous in to later years.

The only randomised controlled clinical trial of hormone replacement therapy and cardiovascular di-

sease reported to date did use combined cyclic estrogen progesteron therapy. In that ten years study, Nachtigal et al (21) found the relative risk of heart attack in the group treated with estrogen progestin was one third of the placebo group.

The other data that supports cardioprotective properties of estrogen replacement is from study of Sullivan et al (42). If the women with angiographically proved CHD were taking estrogen, the overall risk of death due to CHD was reduced by 60% when the other factors such as smoking and cholesterol levels were taken into account. If severity of coronary occlusion is assessed by angiography another aspect of replacement can be evaluated. Women taking estrogen had change of 60% lower risk for having moderate or severe occlusion than did women who were not taking estrogen (43). Whether women were treated with 0.625 or 1.25mg. conjugated estrogen there was 50% reduction in mortality due to myocardial infarction compared with untreated women. As a result if the goal is maintenance of longevity, reduction in CVD spare 90% of the total lives saved in estrogen replacement therapy. Prevention of CVD alone by estrogen replacement cause an overall decrease in mortality 5250 per 100,000.

All of the above epidemiologic data pertain to the use of estrogen alone. Addition of progesteron which is the major complement of estrogen replacement in gynecologically intact women should also be evaluated. Although addition of progesteron seems to attenuate the effect of estrogen onto cardiovascular risk,

Table 1. Summary of studies of replacement of estrogen and cardiovascular disease

Study	Year	Population	End points	Relative risk	P value
Nachtigal (21)	1979	84	Fatal/Nonfatal "MI"	0.33	p>0.05
Talbot (22)	1977	64	Suddenddeath	0.34	p>0.05
Ross (23)	1981	133	Fatal "CHD"	0.43	p<0.01
Sziko (24)	1984	36	Nonfatal "MI"	0.61	p>0.05
Adam (25)	1981	76	Fatal "MI"	0.65	p>0.05
Pfeffer (26)	1978	185	Fatal/Nonfatal "MI"	0.68	p>0.05
Rosenberg (27)	1976	336	Fatal/Nonfatal "MI"	0.97	p>0.05
		6730 controls			
Rosenberg (28)	1980	477	Nonfatal "MI"	1.00	p>0.05
		1832 controls			
Jtak (29)	17	17	Nonfatal "MI"	7.5	p>0.05
Lafferty (30)	1985	124	Fatal/Nonfatal "MI"	0.16	p=0.05
Macmahan (31)	1978	1891	All "CVD"	0.30	NA
Stampfer (32)	1985	32317	All "CVD"	0.30	p<0.01
Hammond (33)	1979	610	All "CVD"	0.33	p<0.01
Potocki (34)	1971	198	All "CVD"	0.33	NA
Bush (35)	1983	2270	"CVD" mortality	0.34	p<0.05
Burch (36)	1974	737	Fatal "CHD"	0.43	p<0.05
Petiti (37)	1979	16638	"CVD" deaths	0.50	p<0.05
Henderson (38)	1986	7610	Fatal/Nonfatal "MI"	0.54	p<0.05
Paganini (39)	1988	8832	Fatal stroke	0.53	p<0.05
Wilson (40)	1985	1234	All "CVD"	1.76	p<0.05

Table 2. Commonly used oral estrogens

Preparation	Dose
Conjugated equine estrogens	0.625 to 1.25mg
Piperazine estrogen sulfate	0.625 to 1.25mg
Micronized 17 beta estradiol	1.0 to 2.0mg
17 alfa ethinyl estradiol	0.01 to 0.02mg

there is still potential benefit to maintain postmenopausal women with this regimen.

Nearly all the studies suggesting protective effect of estrogen replacement therapy relate to use of unopposed oral estrogens, and relatively small number of studies can be found to determine the effects of long term combined therapy. However, one study reported that the addition of a progesterone does not exert opposite effects (45).

Reduced level of HDL cholesterol is largely caused by androgenic properties of the progesterone. As a rule it is expected that the added doses of progesterone results in a fall in HDL cholesterol, and an increase in LDL cholesterol levels (46). Both C19 nortestosterone (noretindrone, norgestrel, norethindrone acetate) and C21 derivatives share that characteristic. But medroxyprogesterone acetate (MPA) reduce HDL cholesterol only slightly at a doses of 10mg. (47). Where as levonorgestrel is the most potent progesterone in the androgenic group, natural progesterone has no major effect on HDL2 cholesterol.

The other negative effect of progesterones is, although this is yet primarily a theoretical concern, a negative influence on vessel wall physiology by counteracting the beneficial changes mediated by estrogen. It has been suggested that even natural progesterone prevents the marked increase in prostacyclin levels induced in human umbilical arteries by estrogen (48).

The gold standard have been well defined in discussions at an international consensus conference held in 1988 (49). In view of the potential negative impact of progesterones on cardiovascular system, imperative rules of administration of progesterones can be summarized as; prescription in the lowest possible effective doses, avoidance of use of the more androgenic compounds and sparing the women who have had a hysterectomy from progesterone.

Based on available data, the daily doses necessary for endometrial secretory transformation in most patients are as follows; MPA 5-10mgr, noretindrone or noretindrone acetate, 0.7-1.0mgr; dl norgestrel, 150 microgr; and micronized oral progesterone 300mgr.

Still further developments and alternative strategies are needed for the optimal estrogen replacement therapy. A new generation of progesterone have being developed; such as desogestrel and gestodene. Des-

ogestrel which minimizes the metabolic impact when prescribed in the contraceptive pill may substitute by more androgenic compounds (50).

An alternative way for avoiding hepatic metabolism is administration of progesterone transdermals in the estrogen containing patch are currently being investigated.

The rationale for the introduction of "continuous / combined therapy" is, production of atrophic endometrium with very small progesterone doses that achieve better lipid impact and to lessen, physiological side effects. However, published data on this kind of administration are relatively sparse. Favorable lipid profile was reported from the study of Hargrove et al (51). This statement is based on a 10% fall in total cholesterol and a 50% increase in HDL cholesterol. The other study was from Mattsson et al (52) who were the first to evaluate the effects of continuous regimen. In two of the four group (norethisterone acetate 0.5mg and medroxyprogesterone acetate 2.5mg) the 10-15% reduction in LDL cholesterol seen at 4 months had almost disappeared at the end of 12 month period. However, whether these changes were due to factors other than treatment is not clear (52).

As a summary there is little basis for confidence that continuous / combined therapy has a more favorable impact on lipid and lipoprotein metabolism than does sequential therapy. A large, prospective, controlled study is required to assess thorough effects of this mode of therapy on to cholesterol metabolism.

TYPES OF ESTROGEN AND ORAL VERSUS TRANSDERMAL ADMINISTRATION

Traditionally, estrogens used for replacement therapy in postmenopausal women have been given orally. Transdermal administration is gaining popularity and has been accepted alternative to oral replacement therapy. Only available form of oral estrogen in Türkiye is conjugated equine estrogens. Estrogen taken orally results in the virtually direct provision of pharmacologic amounts of biologically potent estrogen to the liver. Then potency of type of estrogen determines hepatic response. Potentially harmful induction of clotting factors can result from the use of large doses of a potent estrogen such as ethinyl estradiol. Natural estrogens, including estradiol and estrone and conjugated equine estrogens have less marked effects than the major synthetic estrogens, ethinyl estradiol and DES. Maschak et al (53) reported that the natural estrogens had no major effect on the induction of globulins, though a slightly greater effect was seen with conjugated equine estrogens than with estrone sulfate and micronized estradiol. Ethinyl estradiol produced an effect that was more than 200 times potent. Natural oral estrogen has no deleterious effect on coagulation. Studies by Note-

lovitz et al (54) have confirmed that the coagulation factors such as factor VII, factor X, and antitrombin III are not altered in recipients of natural estrogen replacement. For this reason, synthetic estrogens should not be used for menopausal treatment unless the doses are extremely small.

Although the hepatic first pass effect produces undesirable conditions with regard to hepatic globulins. It also means of increasing HDL cholesterol. Contrary to current belief, this effect is not specific to oral administration. Estrogen via non oral routes in sufficient doses and for a sufficient time has been found to exert a beneficial effect on lipoprotein profiles. Lobo et al (55) showed that transdermal estradiol (0.1 mg) had an effect on total cholesterol and HDL cholesterol comparable to that of subdermal pellet (50mg) after 6 months of therapy.

Recently we have demonstrated that transdermal route is effective as oral estrogen therapy in maintaining favorable lipid profile in women who underwent total abdominal hysterectomy and bilateral salpingoophorectomy (unpublished data presented in Third International Congress of OB & GYN in Izmir).

As a conclusion; almost all epidemiologic studies of postmenopausal hormone replacement therapy with unopposed estrogen have found a reduced risk of death and cardiovascular disease in treated women. Since progestins have varying degrees of androgenic activity, combined estrogen progestin regimens may not be associated with similar protection. Clinical trials are needed to examine the relative risk and benefit of different hormone regimens in postmenopausal women.

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