The action of ovarian hormones in cardiovascular disease

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ABSTRACT

The incidence of cardiovascular disease (CAD) differs between men and women, in part because of differences in risk factors and hormones. This sexual dimorphism means a lower incidence in atherosclerotic diseases in premenopausal women, which subsequently rises in postmenopausal women to eventually equal that of men. These observations point towards estrogen and progesterone playing a lifetime protective role against CAD in women. As exogenous estrogen and estrogen plus progesterone preparations produce significant reductions in low-density lipoprotein (LDL) cholesterol levels and significant increases in high-density lipoprotein (HDL) cholesterol, this should in theory lower the risk of CAD. However, results from oral contraceptive (OC) use and combined estrogen and progesterone hormone replacement therapy (HRT) have suggested that hormone replacement regimes do not provide cardiovascular protection. In fact, depending on the preparation and the presence or absence of genetic risk factors, an increased risk of cardiovascular diseases such as venous thrombosis, myocardial infarction (MI) and stroke have been observed. Interestingly, in the majority of studies the increase in risk was highest in the first year, after which an increase in risk was not observed, and in some studies a lower risk of CAD was evident after four or five years of exogenous hormone administration. While the debate continues about the merits of HRT, and several good reviews exist on the statistics of CAD in relation to exogenous hormones, we have decided to review the literature to piece together the physiological actions of estrogen and progesterone preparations on the individual mechanistic components leading to CAD; namely, the altered endothelium and the haemostatic balance between coagulation and fibrinolysis. We present possible mechanisms for how HRT and OCs protect against MI in the absence of cardiovascular risk factors but increase the incidence of MI in their presence. We also speculate on the roles played by hormones on the short- and long-term risks of cardiovascular disease.

Key terms: hormone replacement therapy (HRT), oral contraceptives, atherosclerosis cardiovascular disease, estrogen, progestins, venous thrombosis, myocardial infarction.

INTRODUCTION

Cardiovascular disease (CAD) is a general description incorporating several pathologies including coronary heart disease, stroke and venous thrombosis. CAD is the main cause of morbidity and mortality in Western countries (Murray & Lopez, 1997). Differences in the incidence of hypertensive heart disease and coronary heart disease, atherosclerosis and cardiac remodeling after myocardial infarction (MI) between males and females are well established (Mendelsohn & Karas, 1999; Stevensen, 2000). Estrogen improves well-defined risk factors, such as lipid profiles; it also has direct effects on the myocardium, endothelium, and vascular smooth muscle (VSM). Estrogens enhance flow of cholesterol from the diet through chylomicrons and chylomicron remnants to the liver, through very low-density lipoprotein (VLDL) and low-density lipoprotein (LDL) to cells, and through reverse cholesterol transport from cells via high-density lipoprotein (HDL) to the liver to be finally eliminated in the bile and intestine (Knopp, 1997). Estrogen increases levels of VLDL and subsequently the levels of triglycerides, decreases LDL levels due to the up-regulation of LDL receptors, and increases HDL due to increased secretion.
of apoA-I and reduced removal of HDL due to a reduction in hepatic lipase activity (Knopp, 1997; Espeland et al., 1998; Zhu et al., 1999; Mendelsohn & Karas, 1999; Table I).

Estrogen is widely regarded as having beneficial effects on the three layers of the arterial wall; the intima (endothelium), the media, and the adventitia. These beneficial effects include reduction in plasma fibrinogen, plasminogen activator inhibitor (PAI-1) activity, reduced LDL oxidation in plasma, enhanced glucose metabolism, and enhanced insulin resistance (Knopp, 1997; Espeland et al., 1998; Mendelsohn & Karas, 1999; Cushman et al., 1999; Sack et al., 1994; Table I). In the arterial endothelium, estrogen increases nitric oxide (NO) synthase activity and NO production (Hishikawa et al., 1995; Caulin-Glaser et al., 1997; Table I). NO is beneficial to arterial vasomotion in women who have angina pectoris due to vasospasm (Guetta and Cannon, 1996). In the intima and media of the arterial wall, estrogen reduces calcification and secretion of inflammatory cytokines, such as fibroblast growth factor (FGF), inter-cellular adhesion molecule (ICAM-1), vascular cell adhesion molecule (VCAM-1), endothelial- and platelet-selectin (E- and P-selectin), nuclear factor kappa B (NFκB) and consequently reduces the atherosclerosis which is associated with release of these cytokines in animal models (Knopp, 1997; Guetta and Cannon, 1996; Adams et al., 1990; Haarbo et al., 1991; Table I).

Conflicting results have been reported on the effects of progestins on atherosclerosis. Several studies demonstrate no effect in both primate and cholesterol-fed rabbit models (Adams et al., 1990; Haarbo et al., 1991). However, other studies have shown that natural progesterone and synthetic progestins oppose the beneficial effects of estrogen (Hanke et al., 1996; Adams et al., 1997). Progestins have been reported to oppose the estrogen-induced increase in plasma NO metabolites (Imthurn et al., 1997) which indicates that progestins inhibit NO production in endothelial cells and is further evidence of the proatherogenic effects of progestins in the presence of estrogen. Progestins appear to reduce the stimulatory effect of estrogens on lipoprotein transport in the bloodstream. For example, VLDL secretion is reduced; remnant removal is impaired, LDL receptors are down-regulated, increasing LDL-cholesterol levels; and HDL levels are reduced in response to increased hepatic lipase activity. Progestins also increase glucose, insulin and fibrinogen plasma levels (Knopp, 1997).

The estrogen receptor (ER) is classically a ligand-dependent nuclear transcription factor. However, recent evidence indicates that non-nuclear liganded ER can also regulate the activity of intracellular second messengers and membrane-associated receptors and signaling complexes (Ho & Liao, 2002). In the cardiovascular system, these non-nuclear signaling pathways mediate rapid vasodilatation (White et al., 1995), inhibition of the response to vessel injury (White et al., 1997; Sullivan et al., 1995), reduction in myocardial injury after infarction (Node et al., 1997), and attenuation of cardiac hypertrophy (Douglas et al., 1998). The importance of estrogen in the cardiovascular system has been elucidated from ER knockout and mutation studies. In a single case study, a young man presented with homozygous disruption of the ERα gene, which resulted in the expression of a truncated receptor lacking both DNA and hormone-binding domains. This patient developed premature CAD and impaired brachial endothelium-dependent vasodilatation providing further evidence for a developmental and protective role for estrogen in the heart and protection from CAD (Sudhir et al., 1997a; Sudhir et al., 1997b). Studies in ovariectomized mice show that 17β-estradiol inhibits intimal and medial vascular smooth muscle proliferation (Sullivan et al., 1995), implying a protective role for estrogen on both endothelial cells (EC) and vascular smooth muscle cells (VSMC). In ER knockout mice, 17β-estradiol also inhibits medial thickening and VSMC proliferation in carotid injury studies (Iafrati et al., 1997). This suggests that the vascular protection produced by estrogen may be mediated in an ERα-independent manner. Furthermore,
when hearts from these mice are subjected to ischemia-reperfusion they present a greater degree of global ischemia and higher incidence of arrhythmias (Zhai et al., 2000). Studies which show that ovariectomized ERαKO mice exposed to cerebral ischemia have strokes which affect a greater area of the brain also demonstrate that ERα mediates the neuroprotective effects of estrogen (Dubal et al., 2001). Evidence is also mounting that ERβ also has a role to play in the cardiovascular system. ERβ expression is induced in VSMC after vascular injury (Linder et al., 1998), and ERβKO mice are hypertensive and have VSMC ion channel dysfunction (Zhu et al., 2002).

OVARIAN HORMONES AND CARDIOVASCULAR DISEASE

Venous disease

Although sexual dimorphism converts to a lower incidence in arterial diseases in premenopausal women, exogenous hormones in the form of OCs and postmenopausal HRT have been associated with increased risk of venous thrombosis, MI and stroke. The presence of hormone preparations appears to add to the increased risk of venous thrombosis caused by genetic factors such as personal or family history, Factor V Leiden, deficiencies of protein C, protein S, or antithrombin III and hyperhomocysteinemia. Emerging studies show that in vitro fertilization treatment and ovulation induction are also risk factors for venous thrombosis (Bloemenkamp et al., 2003). The risk for venous thrombosis is highest during the first year of OC use. Furthermore, OC users with inherited clotting defects develop venous thrombosis, not only more often, but also sooner, than do those without inherited clotting defects (Bloemenkamp et al., 2000). The nature of the exogenous hormone regime is also a risk factor (WHO, 1995). While the concentrations of estrogen have been declining since the first OC usage in the late 1950s, from values as high as 100 ug mestranol to 15 ug ethinyl estradiol, the progestin dosage has remained relatively constant. The World Health Organization (WHO) concluded that although current users of estrogen and progesterone combined contraceptives have a low absolute risk of venous thromboembolism, their risk is still three to six times greater than that of nonusers, with the risk probably being highest during the first year of use (WHO, 1997). Although conflicting reports exist, the increase in venous thrombosis risk remains constant despite changes in the nature and dosage of estrogen. The progestin component, essential to suppress ovulation, has changed only in composition from first generation to third generation. As with estrogenic compounds, each progestin has been associated with an increased risk of CAD. This risk has been demonstrated to be greater for the newer third-generation than from second-generation progestins (Farmer & Lawrenson, 1998).

The picture for CAD is no brighter in women taking HRT preparations. Unopposed estrogen treatment, while delivering beneficial relief from postmenopausal symptoms, also results in endometrial disorders such as endometriosis and cancer (Smith et al., 1975; Berger & Fowler, 1997). Estrogen replacement therapy is still given to women who do not have a uterus, and medroxyprogesterone acetate (MPA) is added to the majority of current HRT preparations to counteract the endometrial abnormalities which arise from unopposed estrogen administration. In a trial of oral conjugated equine estrogen plus MPA, no overall reduction in CAD events was observed in postmenopausal women with established coronary disease. However, the treatment did lead to an increase in the rate of thromboembolic events and gallbladder disease (Hulley et al., 1998). As with OC, the intriguing observation was made that the risk of thrombotic events was higher in the first year of use, and there was a suggestion that the risk decreased to below control (placebo) levels after four to five years of treatment (reviewed in Rosendaal et al., 2002). This emerging pattern for exogenous hormone preparations indicates
that there is a decreased risk of arterial disease while at the same time an increased risk for venous thrombosis. Whether a common mechanism of coagulation and inflammation contributes to both responses is unclear. Furthermore, there are two separate phases in venous thrombosis: a significant increase in risk in the short term, with a plausible reduction in risk associated with long-term use.

Arterial disease

Atherosclerosis is a progressive disease which is characterized by the accumulation of lipids and fibrous elements in the walls of large arteries and constitutes the most important factor in the growing incidence of CAD. Several risk factors, such as cigarette smoking, diabetes, hypertension and elevated serum lipid concentration, have been shown to increase the incidence and accelerate the progression of the disease (Multiple risk factor intervention trial, 1982; Castelli, 1996). Atherosclerosis is responsible for MI and stroke, as well as for their respective precursor disorders, angina pectoris and transient ischemic cerebral attacks (Born et al., 1991). Atherosclerosis is a focal intimal disease of arteries from the aorta down to vessels of 3 mm external diameter. However, not all arteries are equally susceptible; the internal mammary artery is mostly unaffected while coronary arteries are at high risk (Davies and Woolf, 1993). The increased risk of MI and ischemic and hemorrhagic stroke associated with oral contraceptive usage appears elevated only in women with hypertension or who smoke, the risk being negligible in the absence of these risk factors.

Clinical symptoms of atherosclerosis depend on four mechanisms: 1) lipid accumulation and connective tissue matrix production can increase plaque volume so that it encroaches on the lumen and impedes blood flow. 2) A plaque can enter an unstable phase and fissure which leads to thrombus formation. The thrombus can encroach or occlude the lumen or, alternatively, embolize, impact and occlude a smaller distal vessel. 3) Although atherosclerosis is a focal disease, it is associated with a generalized abnormality in vascular tone in affected vessels which favors vasoconstriction, especially during stress and exercise. 4) Medial atrophy and destruction can lead to aneurysm formation (Davies and Woolf, 1993).

Many hypothesis of atherogenesis have been proposed. These tend not to be mutually exclusive and differ more in the emphasis given to particular events rather than to opposing points of view. It is well accepted that lipid accumulation in the arterial wall, caused by hyperlipidemia, is the initial step. Recently, advances in cellular and molecular biology have focused on the role of inflammation in atherogenesis (Figure 1). In many animal models, signs of inflammation are observed simultaneously with lipid accumulation. In both experimental animals and humans, blood leukocytes, mediators of the immune response and inflammation, attach to the endothelial cells that line the intima. The normal endothelium does not support the binding of white blood cells. However, soon after the start of an atherogenic diet, endothelial cells begin to express on their surface adhesion molecules capable of binding leukocytes (Huo & Ley, 2001). Among these adhesion molecules, VCAM-1 and ICAM-1 bind monocytes and T lymphocytes found in early atheroma (Steps 1 & 2, figure 1) (Libby et al., 2002a, b). Increased VCAM-1 expression is also localized to sites prone to atherogenesis, such as branch points in arteries where endothelial cells are subject to disturbed blood flow (Topper et al., 1996). Serum levels of soluble P-selectin and ICAM-1 are also elevated in patients with peripheral atherosclerotic disease (Huo and Ley, 2001). Abnormal laminar shear stress reduces NO production which in turn increases VCAM-1 expression (De Caterina et al., 1995). Progestins inhibit NO production (Imthurn et al., 1997). It is widely reported that estrogen replacement therapy decreases VCAM-1 expression, thereby providing a potentially-protective mechanism to early atherogenic processes (Nathan & Chaudhuri, 1997; Seljeflot et
Figure 1. Hormonal effects on the atherogenic process. A. expression of inflammatory and adhesion molecules; initial lipid infiltration and accumulation. B. Plaque growth and increased LDL deposition. C. Plaque rupture and thrombus formation.
An increase in reactive oxygen species (ROS) in the endothelium, intima and adventitia also plays a major role in the deposition of LDL. Risk factors, such as hyperhomocysteinemia and smoking, increase the oxidation of LDL (oxLDL) and thus deposition through expression of scavenger receptors in the arterial wall (Harrison et al., 2003). Increased shear stress may also induce formation by VSMC of proteoglycans that bind lipoprotein particles, facilitating their oxidative modification and further inducing an inflammatory response (Lee et al., 2001) (Step 3, figure 1). Once the leukocytes have adhered to the endothelium and an inflammatory response is initiated, monocytes penetrate the intima in response to monocyte chemoattractant protein-1 (MCP-1) (Gu et al., 1998), (Step 4 & 5, figure 1). Once in the arterial wall, the monocytes differentiate into macrophages in response to macrophage colony-stimulating factor (M-CSF) (Step 4 & 5, figure 1) (Qiao et al., 1997). Estrogen decreases the expression of MCP-1 and M-CSF thereby decreasing monocyte adhesion to endothelial cells and monocyte migration into the subendothelial space (Nathan & Chaudhuri, 1997). Monocyte adhesion and migration produce a localized inflammatory response in which tumor necrosis factor beta (TNF8) and interferon gamma (INFγ) are released by macrophages (Step 6, figure 1). During this stage, expression of C-reactive protein (CRP), a marker of inflammation, is also increased. Combined estrogen and progesterone therapy increases CRP expression and therefore may further increase the damage induced by the inflammatory response (van Baal et al., 1999b, c; Skouby et al., 2002). These macrophages also have increased expression of the scavenger receptor A and CD36 which internalize modified lipoproteins (minimally modified and oxLDL), accumulating cholesteryl esters in the form of cytoplasmic droplets, leading to foam cell formation which characterize the fatty streak, a hallmark of the early atherosclerotic lesion (Step 7 & 8, figure 1) (Libby, 2002a, b). Estrogen also decreases LDL oxidation, thereby preventing foam cell formation and lesion progression (Crook, 2001; Tedeschi-Reiner & Reiner, 2001; Table I).

Growth factors, inflammatory mediators and proteolytic enzymes released by foam cells induce smooth muscle cell replication, which accumulate in the plaque and lay down extracellular matrix, transforming the fatty streaks into a complicated atheroma. As the lesion grows, it narrows the arterial lumen, interfering with blood-flow and causing clinical symptoms, such as angina pectoris or acute MI (Steps 9-11, figure 1). This apparently smooth progression of plaque growth is frequently marked by bursts in growth of the atheroma (Yokoya et al., 1999). There is evidence to suggest that physical disruption of plaques may trigger thrombosis and promote sudden

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expansion of the atheroma (Davies, 1996). The most common mechanism of plaque disruption is a fracture of the plaque’s fibrous cap through the elaboration of proteases, such as the matrix-degrading neutral metalloproteinases. This cap serves to separate the thrombogenic lipid-rich core of the atheroma from the bloodstream, which contains coagulation factors. Fissure of the fibrous cap allows contact between coagulation factors and tissue factor (TF), the main pro-thrombotic element in the lipid core, causing blood coagulation. Subsequently, platelets become activated by thrombin, generated by the coagulation cascade and by contact with the intimal compartment, causing thrombus formation. If the thrombus occludes the artery, it can cause an acute MI (Libby, 2002a, b).

HORMONES AND THE COAGULATION PATHWAY

The intrinsic coagulation pathway starts with injury to the vessel wall, with exposure of sub-endothelial collagen. The first step in this pathway is the conversion of Factor XII to Factor XIIa. The presence of OCs increases the concentration of Factor XII (Gordon et al., 1983). Factor XIIa promotes the activation of Factor XI to Factor Xla, which in turn activates Factor IX. Factor IX has been shown to be increased in women taking HRT (Lowe et al., 2001). HRT preparations have also been associated with a decrease in plasma levels of Factor VIII (Acs et al., 2002). The combination of Factor IXa and Factor VIII promote the activation of Factor X. On the other side, the extrinsic pathway is initiated by the exposure of Tissue Factor (TF) to circulating Factor VII. TF has been extensively reported to be regulated by exogenous ovarian hormone treatments. Holschermann et al. (1999) reported an increase in TF expression by blood monocytes in the presence of OCs that may favor intravascular clotting activation. Furthermore, several HRT preparations have been demonstrated to lower TF levels in the endothelium, but at the same time increase TF/Factor VIIa activity (Koh et al., 2001). The circulating levels of Factor VII are also altered in response to HRT. In short-term studies (up to one year), cyclic regimes of estrogen and progesterone demonstrate that unopposed estrogen increases Factor VII concentration, an effect which was reversed upon addition of progesterone (Bladbjerg et al., 2002; Lowe et al., 2001; Cushman et al., 1999). Other reports have observed a decrease in Factor VII levels after six weeks of combined treatment HRT (Peverill et al., 2001), while others have shown a differing increase in Factor VII depending on the generation of progestin used in OCs (Kluft 2000). The active complex of TF/Factor VIIa is inhibited by tissue factor pathway inhibitor (TFPI) (Badimon et al., 1999). This protein is down-regulated by estrogen and HRT and OC combinations (Bladbjerg et al., 2002; Peverill et al., 2001; Luyer et al., 2001; Harris et al., 1999), leading to the presence of a hypercoagulable state. Proteolytic cleavage resulting in the formation of Factor VIIa leads, in collaboration with TF, to the activation of Factor X (Factor Xa). This conversion point of the intrinsic and extrinsic pathways promotes the conversion of the soluble blood protein prothrombrin to thrombin. Factor Va also plays a regulatory role in this process. The presence of a mutation in factor V, known as Factor V Leiden, greatly increases the risk of venous thrombosis associated with OCs 30-50 times (Bauer, 2002; Vandenbroucke et al., 2001). Increases in circulating levels of prothrombrin are observed with unopposed estrogen and HRT treatment (Vahkavaara et al., 2001; Peverill et al., 2001, respectively). The net results of these regulations have led to reports of increased levels of thrombin in the presence of estrogen and HRT (Norris et al., 2002; Vahkavaara et al., 2001; Peverill et al., 2001). The amino acid fragment (F1+2), which is generated during prothrombin activation and thus provides a good indication of thrombin levels, was shown to be increased in the presence of HRT, and these levels were higher in women who subsequently developed recurrent venous thrombosis (Hoibraaten et al., 2001; Cano & Van Baal, 2001). Thrombin has several roles; firstly to promote the formation of fibrin from fibrinogen; and secondly, to form a complex with thrombomodulin. This
complex can also down-regulate the circulating levels of thrombin (dashed line with negative sign in Figure 2). In contrast to the above mentioned hormonal regulation of the coagulation cascades, combined estrogen and progestin treatments display anticoagulant activity by lowering the circulating levels of the fibrin precursor fibrinogen (Acs et al., 2002; Cushman et al., 1999; van Baal et al., 1999b; Norris et al., 2002; Table I). HRT has also been reported to increase fibrin turnover (Sidelmann et al., 2003). Finally, in clot formation Factor XIIIa, a transglutaminase, crosslinks the fibrin monomers. The degradation of the clot in the processes leading to fibrinolysis is regulated by sex steroid hormones (Table I). As mentioned above, thrombin forms a complex with thrombomodulin which in turn activates Protein C. Reports are contradictory on the effect of hormones on the levels of activated protein C (APC). Some reports demonstrate higher levels in the presence of HRT (Lowe et al., 2001), while others show a reduction (Hoibraaten et al., 2000; Lowe et al., 2001). APC inactivates Factor V, Factor VIIIa (by proteolytic cleavage) and PAI-1. Interestingly, the risk factor, Factor V Leiden, has diminished sensitivity to APC favoring a coagulation state. In accordance with this effect, elevated Factor VIII levels are associated with an increased risk of deep vein thrombosis (Koster et al., 1995; Kamphuisen et al., 2001). A further anticoagulant action of hormones is demonstrated by both unopposed estrogen (Vahkavaara et al., 2001) and HRT-reducing circulating levels of PAI-1 (Lowe et al., 2001; Koh, 2002; Cushman et al., 1999; Table I). PAI-1 inactivates tissue plasminogen activator (t-PA) and plasminogen activator type-urokinase (u-PA). The conversion of plasminogen to plasmin, which participates in fibrinolysis, is regulated by both u-PA and by t-PA. Interestingly, at this stage, hormones create a balance between promoting and reducing plasmin levels. Levels of plasminogen are increased by both estrogen (Luyer et al., 2001) and HRT (Acs et al., 2002), while t-PA levels have been reported to be reduced by estrogen (Vahkavaara et al., 2001) and HRT (Lowe et al., 2001; Koh, 2002). However, Hoetzer et al. (2003) reported that estrogen could increase t-PA levels 30% compared to controls, while the addition of progesterone, in HRT, reversed this effect. As it is clear that hormones have a strong procoagulative role (on the generation of thrombin) and that there exists a fine hormonal balance in the promotion or inhibition of fibrinolysis, depending on the regime and dosage, hormones may increase or decrease the coagulative potential and thus impose either positive or negative effects on cardiovascular disease.

COAGULATION AND CAD

Coagulation and venous thrombosis

Estrogen and combined estrogen and progestin exogenous preparations alter the coagulable state (reviewed in Figure 2). An increase in the concentration of blood procoagulants may initiate clotting and lead to thrombosis formation increasing the risk of venous thrombosis. This clot may disengage from the vessel wall and flow, via the right side of the heart, to the pulmonary arteries causing pulmonary embolism. The increase in venous thrombosis risk is concentrated in the first year of treatment. As demonstrated in Figure 2 and the accompanying text, exogenous hormones maintain a delicate balance between pro- and anti-coagulative states. To date there is insufficient clinical data relating to the levels of coagulation cascade intermediates to draw firm conclusions, but initial reports suggest there may be an increase in fibrinolytic activity with duration of hormonal usage (Salobir et al., 2002).

Coagulation and atherosclerosis

Clotting at the site of the lesion may involve both the intrinsic and extrinsic coagulation pathways (Khrenov et al., 2002a). The extrinsic, TF-dependent, pathway plays a major role in determining the thrombogenicity of atherosclerotic lesions.
and subsequent generation of acute coronary syndromes (Toschi et al., 1997). In the normal vessel, TF is not expressed on endothelial cells or monocytes which are exposed to the circulating blood. TF is restricted to the adventitia and VSMC of the media (Wilcox et al., 1989). However, in atherosclerotic lesions, high levels of TF mRNA and protein can be found in all three cell types which make up the vessel wall (Moreno et al., 1996). Furthermore, TF expression levels have a high degree of correlation with plaque severity and vulnerability (Ardissino et al., 1997).

Although the extrinsic pathway is responsible for the initiation of thrombus formation, there is mounting evidence that it is solely responsible for the occlusive thrombus formation (Sramek et al., 2001; Bilora et al., 2001; Rosendaal et al., 1990; Triemstra et al., 1995). This implies that the intrinsic pathway, which activates factor X 50-fold more efficiently and amplifies the coagulation triggered by the TF-dependent pathway (Mann, 1999), also contributes to thrombogenicity of the atherosclerotic lesion.

Recent evidence indicates that the LDL/HDL ratio may also affect coagulation. Increased LDL concentrations have a procoagulant effect (Moyer et al., 1998) whereas HDL acts as an anticoagulant (Griffin et al., 1999). The mechanism responsible for these effects seems to be lipoproteins providing a phospholipid surface where the assembly of enzymatic complexes of the coagulation cascade can take place. Specifically, VLDL and oxLDL can support prothrombinase activity and LDL supports extrinsic and intrinsic Xase activity (Khrenov et al., 2002b). While the ability of lipoproteins to support coagulation complex assembly is far less than that observed in platelets, enriched LDL and oxLDL present in the lipid-rich core may play an important role in Xase

**Figure 2.** Hormonal effects on the coagulation cascade.
and prothrombinase complex assembly, adding to the thrombogenicity of the lesion. As mentioned above, estrogen decreases LDL and increases HDL levels, while progestins have the opposite effect.

The accumulation of oxLDL within the plaque induces pathological changes in all vessel wall cells, increasing thrombogenesis (Khrenov et al., 2002a). Oxidized LDL induces TF expression in endothelial cells (Fei et al., 1993), macrophages (Brand et al., 1994) and VSMC (Penn et al., 2000). Accumulated oxLDL may also enhance the ability of atherosclerotic lesion cells to form the phospholipid surface required for the assembly and activity of enzymatic complexes of the intrinsic pathway. Exposure of human macrophages and VSMC to oxLDL increases their ability to support Xase and prothrombinase complex activity, greatly increasing thrombin formation (Ananyeva et al., 2002). This increase in intrinsic pathway procoagulant activity is related to increased expression of factor VIII binding sites and more efficient assembly of Xase complex due to increased exposure of phosphatidylserine (PS) on oxLDL-treated cells (Wintergerst et al., 2000). This data indicates that the intrinsic pathway may play an important role in upregulating the thrombogenicity of atherosclerotic lesions following endothelial layer removal and subsequent exposure of VSMC and macrophages to blood flow.

Although apoptotic cells are absent from normal arteries, they are common in advanced plaques and include VSMC, macrophages and T lymphocytes (Bjorkerud & Bjorkerud, 1996). Apoptosis can be induced by a variety of agents; however, apoptosis is probably mediated by oxLDL within the plaque (Okura et al., 2000). One of the characteristics of apoptotic cells is the translocation of PS from the inner to the outer surface of the plasma membrane by a loss of membrane phospholipid asymmetry. The increased expression and accessibility of PS on the outer surface of the plasma membrane increases the thrombogenic potential of these cells by providing a platform for the assembly of complexes of both intrinsic and extrinsic coagulation pathways (Gilbert & Arena, 1996). Another thrombogenic effect of apoptotic cells present in the plaque is the shedding of PS- and TF-rich microparticles (Mallat et al., 1999; Bombeli et al., 1997).

Thrombogenicity is related to up-regulation of both intrinsic and extrinsic pathways. However, down-regulation of anticoagulant and fibrinolytic activity may also contribute to atherothrombosis (Khrenov et al., 2002a). Oral contraceptives have been demonstrated to lower the sensitivity of factors involved in thrombin generation to APC, thus promoting a more coagulative state (reviewed in Cano & Van Baal, 2001). There is further evidence which shows that thrombomodulin and the endothelial cell protein C receptor expression are down-regulated in atherosclerotic plaques (Laszik et al., 1994). Since HDL increases protein C activity, this down-regulation may be associated with decreased HDL levels observed in atherosclerosis (Griffin et al., 1999).

**THE BENEFITS AND THE RISKS OF EXOGENOUS HORMONES IN ARTERIAL DISEASE**

The paradox of why exogenous hormones increase the short-term risk of cardiovascular disease yet may lower long-term evident risk may be related to the nature of the genesis of venous thrombosis and MI. MI is a consequence of atherosclerosis, while venous thrombosis is a disease of the veins which arises from hypercoagulation and venous stasis. Arteries are larger than veins and have greater flexibility brought about presence of an elastic layer. Movement of the wall of the endothelium is maintained by a delicate balance of dilating factors, such as NO and bradykinin, in combination with constricting factors, such as angiotensin-II, thromboxane, and endothelin among others (reviewed in Cano & Van Baal, 2001). These may be contributing factors as to why arterial disease and myocardial infarction have a lower incidence than venous thrombosis. Another important
Figure 3. Role of hormone replacement therapy (HRT) in atherogenesis in women with and without cardiovascular risk factors.
factor is the process of atherogenesis which occurs only in the arteries. As previously mentioned, exogenous hormones only increase the risk of MI in the presence of cardiovascular risk factors. As depicted in Figure 3 (Scenario A) using HRT as an example in women with no risk factors the increase in blood procoagulants (hypercoagulative state) will not lead to thrombosis formation, due to the size and movement of the artery, and thus the beneficial effects of estrogen in preventing the formation of an atherosclerotic plaque may provide long-term protection against MI and stroke. These beneficial effects of exogenous hormones include: 1) lowering LDL and raising HDL (Table I); 2) reducing intimal damage on vessel walls (Table I); 3) lowering the expression of adhesion molecules such as E-selectin and sICAM-1 (Van Baal et al., 1999a; Table I); 4) lowering MCP-1 (Stork et al., 2002) and (5) increasing anticoagulant APC activity.

However, when women combine exogenous hormones with previous cardiovascular risk factors, an increase in risk of MI is observed (Figure 3, Scenario B). In this case, a risk factor such as hypertension facilitates incorporation of LDL into the arterial wall (Medina et al., 1997), the first step in the formation of an atherosclerotic plaque. This step is further confounded by smoking, which leads to the oxidation of LDL and thus enhanced deposition. Once an atherosclerotic plaque is formed (or is already present), the lowering by estrogens and/or HRT of soluble ICAM-1 and E-selectin and the increasing levels of CRP and matrix metalloproteinase-9 (MMP-9) will destabilize the plaque (Cano & Van Baal, 2001; Stork et al., 2002; Piercy et al., 2002; Zanger et al., 2000). When rupture of the atherosclerotic plaque occurs, a process which is also promoted by hypertension, the coagulation cascade is initiated. In this instant, the increased presence of TF, and other factors, that exist in the presence of exogenous hormones will promote coagulation and increase the clotting potential, leading to more rapid and greater clot formation and thus thrombosis and MI (Figure 3). If thromboembolism, occurs a high change of stroke will ensue. In support of a hypercoagulable state being a risk factor only in the presence of an atherosclerotic plaque, which is deposited more frequently in the presence of the above-mentioned risk factors, markers of coagulation and fibrinolysis (such as t-PA, PAI-1 fibrinogen, and D-dimer) are not associated with increased risk of myocardial infarction, but are associated with atherosclerosis (Haverkate, 2002).

THE FUTURE OF EXOGENOUS HORMONES

The small increase in CAD in women using OCs does not outweigh the benefits from avoiding the trauma and complications arising from unwanted pregnancies, especially in non-smokers with no prior history or cardiovascular risk factors. Along with beneficial effects on mood, osteoporosis and hot flushes, the HRT preparations were to be a simple and safe prophylactic for heart disease. This idea is based on apparently foolproof logic: premenopausal women have lower CAD, take away the hormones at menopause and the protection is lost, add back the hormones and we get back the protection. What may have appeared to be simple on paper has proved to be a nightmare in the clinic. Women are now facing the scenario that not only are their HRT preparations not delivering a protective effect, but that these preparations may actually be putting them at higher risk for CAD (certainly in the short term).

Unfortunately, there are a number of variables queuing up for consideration. First, we have no evidence that if women never went through menopause, that they would maintain the cardiovascular protection as they grow older, since changes in the coagulation system are known to occur naturally with the aging process. Second, although women develop CAD about ten years later than men, they are likely to fare worse after a heart attack (Giardina, 2000). Third, the hormone regimes given as HRT can never exactly mimic the circulating balance and concentrations present in premenopausal
Neither will exogenous hormones emulate the specificity of *in vivo* action derived from the local expression of hormones.

Although an *in vivo* premenopausal situation is never truly possible, the future of HRT is not dead. The beneficial effects of exogenous hormones are required by a subset of women, and as life expectancy increases with every generation, the postmenopausal phase will account for an ever-increasing portion of a woman’s life.

The negative results from the Women’s Health Initiative study (Wassertheil-Smoller *et al.*, 2003) may reflect the combination of risk factors, such as smoking, with hormonal preparations. In the Women’s Health Initiative trial, 50% of the women on HRT had smoked before or continued to smoke during the study (Mueck & Seeger, 2003). Estrogen turnover is increased in women who smoke, reducing the beneficial estrogenic actions (Mueck & Seeger, 2003).

Through the vast array of data regarding OCs, HRT and cardiovascular disease, a beneficial effect or at the very least an insurance of no increased CAD risk, does appear to be present under tightly-defined parameters. It is evident from these clinical studies that the choice of exogenous hormone, combined with personal history and the presence of cardiovascular risk factors need to be taken into consideration before prescription. The feasibility of screening for CAD risk factors may have to be considered. Although the results are not fully clear on the effects of HRT on CAD in healthy, risk-factor-free women, the data does suggest that HRT should not be prescribed, at present, for the prevention of cardiovascular disease. In regard to preparation and dosage, Rosendaal et al. (2002) recommend that contraceptives with 30 ug ethinyl estradiol should be the first choice and that third-generation progestins should be avoided due to their association with increased venous thrombosis.

The information accumulated to date has mainly concentrated on estrogens as a major risk factor, but increasing evidence supports a role for progestins in pathogenesis. The objectives now facing the scientist and the clinician is to better understand the workings, at the physiological and molecular level, of estrogen and progesterone and to determine where and when hormones are required and apply them accordingly. Hopefully, the foolproof plan was merely naive in its execution, while the logic is still firmly in place.

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