In Vivo and In Vitro Evidences of Dehydroepiandrosterone Protective Role on the Cardiovascular System

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Context: Dehydroepiandrosterone (DHEA) and its sulfate ester, Dehydroepiandrosterone Sulfate (DHEA-S) have been considered as putative anti-aging hormones for many years. Indeed, while DHEAS is the most abundant circulating hormone, its concentration is markedly decreased upon aging and early epidemiologic trials have revealed a strong inverse correlation between the hormone concentrations and the occurrence of several dysfunctions frequently encountered in the elderly. Naturally, hormonal supplementation has been rapidly suggested to prevent DHEA (S) deficiency and therefore, age-related development of these pathologies, using the same strategy as estrogen replacement therapy proposed in postmenopausal women.

Evidence Acquisition: All references were searched using PubMed and the following strategy: our initial selection included all articles in English and we sorted them with the following keywords: "DHEA or DHEA-S" and "heart or vascular or endothelium or cardiovascular disease". The search was limited to neither the publication date nor specific journals. The final selection was made according to the relevance of the article content with the aims of the review. According to these criteria, fewer than 10% of the articles retrieved at the first step were discarded.

Results: In this short review, we have focused on the cardiovascular action of DHEA. We started by analyzing evidences in favor of a strong inverse association between DHEA (S) levels and the cardiovascular risk as demonstrated in multiple observational epidemiologic studies for several decades. Then we discussed the different trials aimed at supplementing DHEA (S), both in animals and human, for preventing cardiovascular diseases and we analyzed the possible reasons for the discrepancy observed among the results of some studies. Finally, we presented putative molecular mechanisms of action for DHEA (S), demonstrated in vitro in different models of vascular and cardiac cells, highlighting the complexity of the involved signaling pathways.

Conclusions: The identification of the beneficial cardiovascular effects of DHEA (S) and a better understanding of the involved mechanisms should be helpful to develop new strategies or pharmacologic approaches for many lethal diseases in Western countries.

Keywords: Dehydroepiandrosterone; Endothelium; Myocytes, Cardiac; Cardiovascular System; Disease; Steroids

1. Context

An excess of steroid hormones such as mineralocorticoids has been linked to heart hypertrophy, arrhythmias, inflammation, fibrosis, and apoptosis as well as to endothelial and smooth muscle vascular dysfunctions. Benefits of antagonizing aldosterone action in patients with cardiovascular diseases (CVD) have demonstrated a causal relationship between steroid hormones and CVD (1). On the other hand, putatively protective steroid hormones such as dehydroepiandrosterone (DHEA) decrease with aging. DHEA, and particularly its sulfated derivative, DHEA sulfate (DHEA-S), are among the most abundant circulating steroid hormones. These two hormones are derived from cholesterol and are principally produced by the adrenal cortex; however, small amounts of them have been proposed to be produced by other organs such as the heart. Since the discovery of DHEA and its sulfated metabolite during the first half of the 20th century, many publications have hypothesized that the development of DHEA deficiencies with age might play a key role in the degradation of many functions and contribute to the genesis of several disorders, including cardiovascular defects (2). Other studies have reported a beneficial role for DHEA (S) in various physiologic or pathophysiologic conditions such as brain development, aging, osteoporosis, immune system-mediated rheumatologic diseases, diabetes mellitus (DM), obesity, chronic heart failure (particularly when linked with oxidative stress), and recently, vascular remodeling as occurring in pulmonary arterial hypertension (3). Nevertheless, only few marked beneficial effects have been clearly reported with DHEA.
supplementation in human and therefore, this subject is still controversial.

In the present article, we aimed at providing a short review of recently published works showing the inverse correlation between the circulating levels of DHEA and DHEA-S and the degree of illness in several pathophysiologic situations. We focused our attention on CVD and on the beneficial effects of counteracting DHEA (S) deficiency by supplementation. We also discussed the putative mechanisms of action exerted by the hormone in several cardiovascular target organs and tissues.

2. Evidence Acquisition

All references have been searched using PubMed and the following strategy:

Our initial selection included all articles in English language and we sorted them with the keywords “DHEA or DHEA-S” and “heart or vascular or endothelium or cardiovascular disease”. The search was not limited by neither the publication date nor specific journals. The final selection was made according to the relevance of the article content to the aims of the review. According to these criteria, <10% of the retrieved articles at the first step was discarded.

3. Results

3.1. Low Levels of Dehydroepiandrosterone (Sulfate) Predicts Cardiovascular Health Disorders

3.1.1. Assessing a Role for Dehydroepiandrosterone (Sulfate)

Correlation between DHEA or DHEA-S plasma levels and ageing-related pathologies has been widely explored during the last twenty years. Indeed, DHEA (S) levels in circulation decrease with ageing more markedly than those of other hormones. Numerous studies have shown a negative correlation between the levels of these hormones and ageing, as defined by the occurrence of several disorders including CVD as well as more definitive entities such as mortality. However, the predictive value is influenced by factors such as sex and smoking habits. In a cohort of elderly subjects, it was demonstrated that DHEAS levels inversely correlated with men mortality whereas there was no association with women mortality (4). In addition, only one published work has clearly mentioned physiologic differences between DHEA and DHEA-S to relate their respective concentrations to various pathologies (5). They highlighted the difficulty to properly correlate DHEA concentration with diseases, probably due to the different hormone distribution throughout life, compared to its sulfated form. The authors also reported that DHEA-S levels are much more often measured in studies than DHEA levels, which are often not reported at all.

3.1.2. Dehydroepiandrosterone (Sulfate) Deficiency and Cardiovascular Risk

The CVD encompass numerous and diverse kinds of pathologies related to peripheral vessels as well as to the heart. Barrett-Connor et al. have demonstrated that mean DHEAS concentrations decreases in men with age and are significantly lower in those with a history of heart disease (6). After adjustment for age and other CVD risk factors, they found that men with lower DHEA-S levels (<3.8 μmol/L) had a 1.5-fold higher risk to die of any cause (which was not statistically significant), but a 3.3-fold higher risk to die of CVD (P < 0.05), and particularly from ischemic heart disease (IHD).

Clinical and demographic parameters are relevant to drawing any conclusion about causal relationship between DHEA (S) and CVD. For example, concerning the sex, studies are more consistent for men, and when looking at women cohorts, the relationship between DHEA-S and mortality from all causes, including from CVD, is often less obvious. Nevertheless, in the WISE (Women’s Ischemia Syndrome Evaluation) study, including 270 postmenopausal women undergoing coronary angiography for suspected ischemia, a positive correlation between DHEAS levels and CVD outcome was reported, which was potentially explained by a better atherosclerosis state of patients with higher DHEA-S levels (7). Another study demonstrated that the decrease of DHEA-S concentrations observed in elderly women is significantly more pronounced in case of coronary heart diseases, osteoporosis or depression (8).

In contrast, prospectively investigating the longitudinal patterns of sex steroids in a cohort of 254 elderly men on a ten-year follow-up did not reveal any significant association between sex steroids (testosterone and DHEA-S) and the incidence of CVD or all-cause mortality risk (9). In conclusion, DHEA-S levels are variably good markers not perfect for predicting CVD in the general population probably because many additional important factors have to be integrated in these predictions.

The role of DHEA and DHEA-S has been studied in the context of endothelial function in postmenopausal women with coronary risks factors. Results have shown that lower levels of DHEA-S are associated with endothelial dysfunction as assessed by a positive correlation between DHEA-S and percentage of flow-mediated dilatation (%FMD), a marker of endothelial function (10). This correlation was independent of other coronary risk factors such as age, body mass index, blood pressure, total and high-density lipoprotein cholesterol (HDL-C), fasting glucose levels, or smoking habits.

Recent studies evaluated whether DHEA (S) is associated with development of carotid atherosclerosis, a situation predicting cardiovascular complications. For example, measurement of intima media thickness (IMT), blood flow volume, and endothelial function (%FMD) has been performed by ultrasonography of carotid arteries in 419...
Japanese patients of both sexes with cardiovascular risk factors, and IMT appeared to be inversely correlated with DHEA-S levels in males, but not in females. In contrast, DHEA-S was a positive factor for predicting blood flow volume in females, but not in males, and no significant correlation could be demonstrated in this study between DHEA-S and %FMD regardless of sex (11).

The association between androgen levels, including DHEA-S and total and bioavailable testosterone, with aortic atherosclerosis has been also investigated in 1032 non-smoking Dutch men and women aged 55 years or older. Intimal atherosclerosis was assessed by radiographic detection of calcified deposits in the abdominal aorta; however, no clear association between levels of DHEA-S and presence of severe aortic atherosclerosis was found in men or women (12).

In order to examine the association between DHEA and coronary atherosclerosis, plasma DHEA and DHEA-S levels had been previously determined in 206 middle-aged patients undergoing coronary angiography. The DHEA-S level was significantly lower in subjects with stenosis and was inversely associated with the number of affected coronary vessels and the extent of coronary atherosclerosis (13). Another part of the same study also investigated the putative protection of DHEA against the development of vasculopathy after cardiac allograft. In a cohort of 61 cardiac allograft recipients, it has been found that DHEA levels inversely correlated with the development of accelerated coronary vasculopathy, which appeared earlier in patients with low levels of the hormone, suggesting that high levels of DHEA could retard the development of coronary atherosclerosis and coronary allograft vasculopathy.

In a prospective study (Massachusetts Male Aging Study) on 167 males aged 40 to 70 years at baseline followed for over ten years, low serum DHEA (S) was predictive of later IHD independent of other risk factors. The subjects in the lowest quartile (DHEA-S < 4.3 μmol/L) were significantly more likely to develop IHD with an adjusted odds ratio of 1.6 (95% CI, 1.07-2.39) (14).

The risk of ischemic stroke might be influenced by DHEA-S levels, as suggested by another study based on a cohort of 32,826 women (Nurses’ Health Study) following patients for cardiovascular events (15). Matching 461 cases of ischemic strokes confirmed by medical records, with controls of the same age, race, menopausal status, hormone use, and smoking status demonstrated that lower DHEA-S levels were associated with a greater risk of stroke.

Association of atrial fibrillation and its prevalence with DHEA-S levels were investigated in 436 men and 544 women aging 65 to 97 years old. Interestingly, a low DHEA-S concentration was significantly associated with cardiac arrhythmias in men, while occlusive arterial diseases, chronic obstructive lung disease, or osteoporosis were more frequent in women with low DHEA-S. However, only the association with osteoporosis remained significant after controlling for differences in lifestyle and general health status parameters (16). The authors concluded that in the elderly, low serum DHEA-S levels are more a nonspecific indicator of aging and health status rather than a risk marker of specific diseases.

Diabetes mellitus is frequently associated with CVD, and significantly lower DHEA-S concentrations were observed in 51 diabetic patients with a specific subtype of type 2 DM in comparison with 49 healthy controls (17). The putative cardioprotective effect of DHEA has been investigated in 62 male patients with type 2 DM by determining the association between the hormone levels and the activated protein C, C-reactive protein, and the IMT of the carotid artery. The study concluded that lower circulating levels of DHEA are associated with decreased activated protein C generation and higher IMT in patients type 2 DM (18).

Finally, testosterone and DHEA were determined in 153 men with DM (age, 65 ± 9 years) with coronary artery disease (CAD). Androgen deficiency, defined as serum levels below the 10th percentile of healthy peers, was common in men with DM with stable CAD and predicted increased cardiovascular mortality (19).

3.2. In Vivo Dehydroepiandrosterone Supplementation: Is It Efficient to Prevent Cardiovascular Diseases?

Several early supplementation studies have been done before any clinical studies aimed at measuring DHEA or DHEA-S levels. This fact led to many misunderstandings as supplementation was not always provided for correcting DHEA (S) deficiencies in patients.

3.2.1. Supplementation in Animal Models

Systemic vascular remodeling, as occurring during restenosis after injury, is characterized by the presence of proliferative and apoptosis-resistant smooth muscle cells with activated Akt pathway. Because DHEA has been previously proposed to inhibit this pathway in cancer, Bonnet et al. investigated a putative role for DHEA in remodeling vessel wall both in vitro, using cultured human carotid vascular smooth muscle cells and saphenous vein grafts challenged with PDGF, and in vivo, with a rat carotid injury model. In each case, DHEA reduced proliferation and increased apoptosis of smooth muscle cells with activated Akt pathway. Because DHEA has been shown to affect the Akt pathway, and DHEA effect was unaffected by androgen or estrogen antagonists, suggesting a steroid receptor-independent mechanism. The authors concluded that orally available DHEA might be attractive for treating systemic vascular remodeling, including restenosis (20).

In order to mimic the metabolic dysregulation often occurring in the elderly, animals are often submitted to a high fat diet. These animals develop inflammatory processes and cardiomyopathies linked to their diabetic state. In this context, DHEA seems to be beneficial for counteracting cardiac disturbances such as unbalanced oxidative status, impairment of cardiac myogenic...
The benefits of DHEA supplementation in the aging population is highly controversial within the literature and therefore, published studies have been recently reviewed in a meta-analysis to determine whether there is a potential therapeutic role of DHEA in postmenopausal women (28). In spite of several observational studies showing that lower DHEA levels were associated with increased cardiovascular risk in women, interventional trials did not demonstrate improvement atherosclerosis or cardiovascular risk factors by DHEA supplementation. The DHEA even appears to be responsible for lowering HDL-C levels, a putative cardioprotective marker, probably through an androgen-dependent mechanism; however, it is not clear how much this action of DHEA is detrimental to the cardiovascular function. The authors conclude that there was no strong current evidence in favor of DHEA supplementation in healthy postmenopausal women.

Several reports also suggest that high doses of DHEA (>25 mg/day) could be potentially pro-arrhythogenic (29) and in most cases, palpitations or tachycardia disappear when the drug is stopped. Whether this deleterious effect of DHEA is related to its in vivo transformation into androgens, or to a direct action of the hormone on cardiomyocytes, remains to be determined.

### 3.3. In Vitro Investigation of Cellular and Molecular Mechanisms of Dehydroepiandrosterone (Sulfate) Cardiovascular Action

When assessing the cellular and molecular mechanism of DHEA (S) action, it is important to realize that because of their quite different chemical properties, DHEA and DHEAS may act differently on their targets cells in terms of both efficiency and specificity of the involved signaling pathways. Moreover, both genomic and non-genomic mechanisms can be recruited by these hormones with divergent consequences for the cell function.

#### 3.3.1. Dehydroepiandrosterone Effect on Isolated Endothelial Cells

Several effects of DHEA (and not always mimicked by DHEA-S) on the endothelium have been extensively described (Table 1). The formation of the atheroma plaque, for example, requires the adhesion of monocytes to vascular endothelium through binding of endothelial neural cell adhesion molecule (NCAM) to monocyte NCAM. This interaction is prevented by sialylation of NCAM. When administered to cultured human coronary artery endothelial cells, DHEA inhibited monocyte adhesion by increasing the expression of polysialylated NCAM, a response sensitive to androgen receptor (AR) and estrogen receptor (ER) antagonists in a sex-dependent manner (30). Interestingly, the expression of polysialylated NCAM was completely abolished in the presence of trilostane, an inhibitor of 3β-hydroxysteroid dehydrogenase, suggesting that DHEA acts via its end metabolites, estradiol and testosterone.
### Table 1. In Vitro or Ex Vivo Assessment of Dehydroepiandrosterone (Sulfate) Mode of Action on Various Cardiovascular Target Tissues

| Target Tissue/Cell (Species) | Macroscopic Biologic Response to DHEA (S) | Receptor Involved (or Messenger) | Cellular Signaling Pathway or Molecular Mechanism Suggested | References |
|-----------------------------|------------------------------------------|----------------------------------|-------------------------------------------------------------|------------|
| **Endothelial cells**       |                                          |                                  |                                                             |            |
| Coronary artery (human)     | Inhibition of monocyte adhesion          | AR and ER (via testosterone or estradiol) | Increased expression of polysialylated NCAM | (30)       |
| Umbilical vein/HUVEC (human) | Inhibition of monocyte adhesion and of TNFα-induced inflammation | N/D | Inhibition of NFκB translocation, reduction of ICAM-1 and E-selectin expression, diminution of ROS production | (31)       |
| Umbilical vein/HUVEC (human) | Prevention of monocyte adhesion induced by high glucose | N/D | Inhibition of NFκB translocation and adhesion molecule expression, antioxidant properties of DHEA (S) | (32)       |
| Aorta (human)               | Inhibition of TNFα-induced inflammation | PPARα | Attenuation of IL-8, ICAM-1 and VCAM-1 expression, inhibition of NFκB translocation | (33)       |
| Umbilical vein/HUVEC (human) | Inhibition of cell proliferation         | NOT through AR or ER (not DHEA-S) | N/D | | (34)       |
| Aorta (bovine) and HUVEC (human) | Increase of cell proliferation and protection against superoxide injury | NOT through AR or ER | Antioxidant properties of DHEA (S), increase of ERK1, 2 and eNOS expression | (26, 35) |
| Aorta (bovine) and HUVEC (human) | Increase of NO release                   | NOT through ER cell surface GPCR | NOS activation through acute (non-genomic) and Pertussis toxin-sensitive pathway | (3, 36)   |
| Aorta (rats)                | Preservation of endothelial function upon aortic stenosis | Sigma-IR | Akt and eNOS phosphorylation and expression, inhibition of Sig-1R downregulation | (37)       |
| **Vascular smooth muscle cells** |                                          |                                  |                                                             |            |
| Pulmonary and aortic artery (rats) | Relaxation of precontracted (KCl) rings | N/D | Opening of potassium channels (cell hyperpolarization) | (38)       |
| Pulmonary artery (rats)     | Reduction of hypoxic pulmonary hypertension, relaxation of artery rings | N/D | Upregulation of the soluble guanylyl cyclase (increased sensitivity to NO) | (39)       |
| Pulmonary artery (rats)     | Relaxation of precontracted (KCl) rings | NOT through AR or ER | Inhibition of T-type calcium channels through a Pertussis toxin-sensitive G protein | (40)       |
| Pulmonary artery (human)    | Inhibition of cell proliferation and sensitization to apoptosis (PAH patients) | N/D | Inhibition of STAT3 and NFAT, induction of mitochondria depolarization and reduction of cell calcium concentration | (41)       |
| **Cardiomyocytes**          |                                          |                                  |                                                             |            |
| Ventricle (rat)             | Reduction of hypertrophy induced by endothelin | N/D | Inhibition of BNP expression | (42)       |
| Ventricle (rats)            | Prevention of corticosteroid-induced hypertrophy and chronotropic response | N/D | Decrease of T-type calcium channel expression and activity, inhibition of ANP and BNP expression; antioxidant properties of DHEA | (43)       |
| Ventricle (rats)            | Improvement of cardiac function and remodeling after chronic hypoxia | N/D | Activation of CREB and PGC1α, increase of myocyte proliferation and respiratory chain activity, reduction of superoxide anions | (24)       |

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*Abbreviations: DHEA (S), dehydroepiandrosterone (sulfate); AR, androgen receptor; ER, estrogen receptor; NCAM, neural cell adhesion molecule; HUVEC, human umbilical vein endothelial cell; TNFα, Tumor necrosis factor; N/D, not determined; NFκB, nuclear factor kappa-light-chain-enhancer of activated B cells; PPARα, Peroxisome proliferator-activated receptor alpha; IL-8, interleukin 8; ICAM-1, intercellular adhesion molecule 1; ROS, reactive oxygen species; VCAM-1, vascular cell adhesion molecule 1; ERK, extracellular signal-regulated kinase; eNOS, endothelial NO synthase; GPCR, G protein-coupled receptor; PAH, pulmonary arterial hypertension; STAT3, signal transducer and activator of transcription 3; NFAT, nuclear factor of activated T cells; and CREB, Response Element Binding Protein.*
The protective action of DHEA against atherosclerosis has been further investigated in human umbilical vein endothelial cells (HUVEC), where the hormone (100 μmol/l) appeared to inhibit in vitro TNFα-induced inflammatory response including NFκB translocation, ICAM-1 and E-selectin expression, reactive oxygen species (ROS) production, and U937 monocyte adhesion (31). The same group previously showed that DHEA was able to inhibit HUVEC proliferation by arresting the cell cycle in the G1 phase, a process that is over-activated in pathologic inflammatory and angiogenic conditions (34). It is noteworthy that the antiproliferative action of DHEA was not inhibited by AR or ER antagonists, was not mimicked by estradiol or testosterone, and more surprisingly, was not mimicked either by the sulfated ester, DHEA-S.

This effect of DHEA on the reduction of endothelial proliferation is in contrast with the observation by others that like estradiol and testosterone, DHEA increased the proliferation of bovine aortic endothelial cells and HUVEC (26). In this case, while the proliferative effect of estradiol and testosterone were abolished by the ER antagonist (ICI 182780) and AR antagonist (flutamide) none of them blocked the proliferative action of DHEA.

The reasons for the existing discrepancy between this finding and the previous study in term of endothelial cell proliferation are not known. High complexity of this system was suggested by an independent observation that testosterone induced the growth of human vascular endothelial cells via AR-independent mechanisms, while dihydrotestosterone (DHT) inhibited the growth of the same cells through an AR-dependent mechanism (35). Moreover, both testosterone and DHT enhanced vascular smooth muscle cell proliferation in a same way and ERK1/2 activation via AR-dependent and AR-independent pathways, respectively.

A protective action of DHEA on human vascular endothelial cells against superoxide injury has been also reported, and this effect appeared to be through AR-independent mechanisms (35), reminding the antioxidant properties of the DHEA molecule. Among other beneficial effects attributed to DHEA, its ability to stimulate nitric oxide (NO) release from endothelial cells is noteworthy. Indeed, DHEA stimulates NO release within five minutes exposure of intact bovine aortic endothelial or HUVEC (36). Increased NO release is due to activation of eNOS and leads to an elevation of cGMP within the cell. Tamoxifen, an ER antagonist, has been shown to block estrogen-stimulated NO release, but not the acute DHEA effect, which was abolished by pertussis toxin. Interestingly, albumin-conjugated DHEA also stimulated NO release, comforting the hypothesis that DHEA exerts its action through a plasma membrane-initiated mechanism, possibly involving a G protein.

Platelets in patients with type 2 DM are more sensitive to aggregation, which has been attributed to a reduced ability of these cells to produce NO. Recently, DHEA was shown to reduce platelet ADP-induced aggregation by 40% in plasma obtained from postmenopausal women with type-2 DM. Using several pharmacologic agents, the authors determined that DHEA action on the platelets was mediated by the activation of the PKCα/eNOS/NO/cGMP pathway (44). Similarly, the increased adhesion of U937 monocytes on HUVEC, induced by high glucose concentrations in vitro, as well as the concomitant overexpression of several adhesion molecules was abolished by DHEA (32). Moreover, DHEA completely blocked the oxidative stress and decreased the translocation of NFκB induced by glucose in these cells, suggesting that DHEA could be useful for preventing some complications of DM and hyperglycemia.

3.3.2. Mechanisms of Dehydroepiandrosterone (Sulfate) Action on Vascular Smooth Muscle Cells

Pulmonary arterial hypertension is a severe disease of the pulmonary vasculature, which is defined by an increased arterial pressure and is characterized by proliferation and resistance to apoptosis of arterial smooth muscle cells. This last phenotype is partly due to the activation by growth factors and inflammatory cytokines of neoplastic pathways involving the transcription factor STAT3 under the control of the protein Src. Similar to cancer cells, Src/STAT3 stimulation leads to the activation of NFAT signaling, itself responsible for both proliferation and resistance to apoptosis.

The DHEA (S) has been reported to be a pulmonary vasodilator that prevent acute hypoxic pulmonary vasoconstriction by opening potassium channels (38) and upregulating the NO-modulated soluble guanylyl cyclase (39), both mechanisms leading to relaxation of vascular smooth muscle cells. Recently, DHEA was shown to reduce the activity of voltage-activated T-type calcium channels within various cell types, including pulmonary artery smooth muscle cells. The modifications of the electrophysiologic properties of the calcium channels observed upon DHEA challenge resulted in the inhibition of KCl-induced contraction of pulmonary artery rings. They were independent of AR and ER and a pertussis toxin-sensitive Gi protein pathway is implicated for these effects (40).

In addition, DHEA reverses human vascular remodeling by inhibiting Src/STAT3 constitutive activation (41). This in vitro inhibition by DHEA was shown to be associated with a reduced activity of NFAT, a mitochondrial membrane depolarization (sensitizing the cell to apoptosis), and a decrease of intracellular calcium concentrations, all these factors contributing to a reduction of cell proliferation. The response to DHEA was similar to that observed upon molecular inhibition of STAT3, suggesting STAT3 as a main target for the hormone in smooth muscle cells.

3.3.3. Specific Effects of Dehydroepiandrosterone on Cardiomyocytes

The putative cardioprotective action of DHEA, which was suggested by several epidemiologic observational
studies, was tested in vitro using a neonatal rat cardiomyocyte culture system. It was found that the presence of DHEA significantly inhibited the increase in myocyte cell size and BNP mRNA levels upregulated by endothelin-1 (42). Interestingly, in the same study, the authors reported that DHEA levels in humans, measured at the coronary sinuses and aortic roots during cardiac catheterization, were suppressed in patients with heart failure in contrast to controls. The opposite was observed for aldosterone concentrations, which were increased in heart failure. They postulated that DHEA or its metabolites exert a cardioprotective action through their anti-hypertrophic effects.

Recently, another study investigated the ability of DHEA to counteract the chronotropic and hypertrophic actions of aldosterone and glucocorticoids (43). Freshly isolated neonate rat cardiomyocytes contract spontaneously in culture and their beating frequency is markedly increased upon stimulation of the mineralocorticoid or the glucocorticoid receptors (MR and GR, respectively). DHEA was found to reduce this chronotropic response by decreasing the T-type calcium channel expression and activity within cardiomyocytes. In the same model, DHEA also prevented MR- and GR-mediated myocyte hypertrophy as well as the expression of ANP and BNP. The kinetics of the negative chronotropic action of DHEA and its sensitivity to actinomycin D revealed the presence of both genomic and non-genomic mechanism of action. While the genomic action was mostly related to MR inhibition, the rapid non-genomic response appeared to be due to DHEA antioxidant properties. The reduction of myocyte hypertrophy and spontaneous beating frequency by DHEA is clearly beneficial for the heart function; however, epiandrosterone, a metabolite of DHEA, has been reported to reduce L-type calcium channel activity in ventricular myocytes and therefore, inhibits myocardial contractility (45).

4. Conclusions

DHEA (S) is the hormone not only circulating at the highest concentration in blood, but it is also displaying the most extensive decrease upon aging. Therefore, it has been tempting to correlate the reduction of the hormone levels with several types of age-related dysfunctions including CVD. Many observational studies performed over decades, demonstrated the existence of a significant inverse association between levels of DHEA (S) and cardiovascular risk, morbidity, and mortality, even after correcting for usual confounding factors. Nevertheless, a strong correlation is not a proof of causality. In order to discriminate between a hormonal deficiency directly responsible for the observed disease or the hormone acting as a simple marker of the biologic age (possibly the main cause of the increased frequency of pathologies), the problem was investigated by measuring the beneficial effects of hormone supplementation. In other words, the question addressed here was whether DHEA (S) was the anti-aging hormone. Unfortunately, supplementation studies were much less conclusive than expected and the benefit of DHEA (S) on the outcome of CVD is still highly controversial. Discrepancies among results can be due to several factors such as insufficient duration of treatment, heterogeneity of the tested population, irreversibility of investigated pathologies, or even economic concerns. Indeed, because DHEA (S) is a cheap, not patentable molecule, already on the free market for a long time in the United States, there is poor motivation in the industry for conducting rigorous and expensive clinical trials.

Therefore, it is not surprising that the strongest information on DHEA (S) action has been obtained from in vitro studies. However, strong limitation in the discovery of the molecular mechanisms of DHEA (S) action are the lack of a specific receptor, which can be clearly identified as responsible for the hormone response, as well as the ability of DHEA (S) to be converted into androgens and estrogens. Moreover, similar to other steroid hormones, DHEA (S) appears to exert its effect through genomic and non-genomic pathways, the latter of which included simple antioxidant properties. Nevertheless, a better comprehension of the basic mechanisms leading to the multiple actions of DHEA (S) on its various target cells will be certainly useful for developing new strategies and better apprehending CVD, which is still killing too many people in Western countries.

Authors’ Contributions

Study concept and design, drafting the manuscript, and critical revision of the manuscript for important intellectual content: Tiphaine Mannic, Joanna Viguie, and Michel Florian Rossier.

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