Abstract

During the last decades, visceral adiposity has been at the forefront of scientific research because of its complex role in the pathogenesis of cardiovascular diseases. Epicardial adipose tissue (EAT) is the visceral lipid compartment between the myocardium and the visceral pericardium. Due to their unobstructed anatomic vicinity, epicardial fat and myocardium are nourished by the same microcirculation. It is widely known that EAT serves as an energy lipid source and thermoregulator for the human heart. In addition to this, epicardial fat exerts highly protective effects since it releases a great variety of anti-inflammatory molecules to the adjacent cardiac muscle. Taking into account the unique properties of human EAT, it is undoubtedly a key factor in cardiac physiology since it facilitates complex heart functions. Under pathological circumstances, however, epicardial fat promotes coronary atherosclerosis in a variety of ways. Therefore, the accurate estimation of epicardial fat thickness and volume could be utilized as an early detecting method and future medication target for coronary artery disease (CAD) elimination. Throughout the years, several therapeutic approaches for dysfunctional human EAT have been proposed. A balanced healthy diet, aerobic and anaerobic physical activity, bariatric surgery, and pharmacological treatment with either traditional or novel antidiabetic and antilipidemic drugs are some of the established medical approaches. In the present article, we review the current knowledge regarding the anatomic and physiological characteristics of epicardial fat. In addition to this, we describe the pathogenic mechanisms which refer to the crosstalk between epicardial fat alteration and coronary arterial atherosclerosis development. Lastly, we present both lifestyle and pharmacological methods as possible treatment options for EAT dysfunction.

Introduction And Background

Coronary artery disease (CAD) is a considerable health condition affecting millions of individuals all over the world [1]. According to the American Heart Association, 840,768 individuals died due to coronary artery disorder in the United States of America (USA) in 2016 [2]. Recent data suggest that CAD remains the principal cause of human mortality and morbidity in both developed and developing countries [3]. Stable angina, acute coronary syndromes (myocardial infarction, unstable angina), and sudden cardiac death are some of CAD’s clinical manifestations [3]. Coronary atherosclerosis, characterized by remodeling and occlusion in the coronary arterial trees, is proposed to be the main etiopathogenic mechanism of CAD [4]. To date, numerous risk factors either modifiable (e.g., arterial hypertension, dyslipidemia, smoking, obesity) or not (e.g., family history of acute coronary syndrome in relatively early adulthood, sex, age) have been associated with the uprising incidence of CAD [5].

In recent years, adipose tissue has been at the center of scientific medical research [6]. A growing body of data has indicated different and crucial roles of human adiposity, from energy fuel to thermoregulation and injury protection [6,7]. In addition to this, data from prestigious experimental and clinical studies have recognized both the endocrine and paracrine effects of adipose tissue on animals and humans [7]. It is now well-established that excessive adiposity releases plenty of inflammatory cytokines leading to a low-grade inflammatory microenvironment, endothelial dysfunction, increase in oxidative stress, and therefore, coronary atherosclerosis [8].

Epicardial adipose tissue (EAT) is the visceral fat unit localized between the myocardium and the inner pericardium, surrounding major coronary vessels [9]. Due to their close anatomic relationship, epicardial fat and the proximal myocardium share the same vessel microcirculation [9]. Numerous clinical and basic research studies have been conducted during the last two decades in order to investigate epicardial fat
To our knowledge, epicardial fat is a biologically active endocrine organ with several paracrine and vasocrine effects on the proximal myocardium [11]. Nowadays, it is accepted that epicardial fat plays a significant and cardioprotective role in normal heart function [8]. However, recent data suggest that abnormal epicardial fat could lead to adverse cardiovascular outcomes through several etiopathogenic mechanisms [12]. In fact, several clinical studies have demonstrated a direct correlation between CAD and dysfunctional EAT [13]. In this way, the unique transcriptome of epicardial fat has become a subject of thorough scientific investigation [9].

In this paper, we briefly report the anatomy, physiology, and quantification methods of EAT. Also discussed are the pathophysiology mechanisms in which abnormal epicardial fat induces CAD. Lastly, we present potential therapeutic approaches in order to modify abnormal EAT's function in CAD.

Review

Anatomy

Epicardial fat is the adipose tissue localized between myocardium and the epicardium (the visceral surface of pericardium), covering approximately 80% of the total heart mass [14]. It is mainly detected in the interventricular and atrioventricular chambers, the apex of the heart and the free wall of the right ventricle surrounding major branches of coronary arterial tree [14]. It is distinguished in myocardial EAT (the adipose tissue just over the myocardial surface) and pericoronary EAT (the fat depot directly located around the coronary arteries) [15]. Epicardial and paracardial fat depots compose pericardial adipose tissue (PAT) [16]. To our knowledge, PAT and epicardial fat share similar morphological features [15]. However, they present different embryological origin, since PAT derives from thoracic mesoderm and EAT from splanchnopleuric mesoderm [16]. Furthermore, EAT is supplied with oxygen and other nutritive ingredients by small branches of coronary vessels, whereas PAT is nourished by thoracic arteries [16]. Histologically, EAT consists of various types of cells, such as adipocytes, inflammatory and immune cells, nerve cells and vascular ones [17]. Additionally, there is no intervening anatomic barrier between myocardium and epicardial fat [18]. As a result, the same small vessel circulation is shared by both of them [18].

Several factors, including age, race, body mass index (BMI) and sex have been associated to anatomical alteration of EAT distribution in the human heart [10]. It has been described that by the age of 65 years EAT thickness increases by almost 22% [19]. Caucasian men are also characterized by greater epicardial fat thickness than black men [20]. Obesity is a triggering factor of increase in visceral adiposity [10]. Thus, abnormal BMI is also correlated to higher EAT mass [10]. Obese women seem to present greater EAT mass than men [10].

Physiology

EAT as an Energy Fat Source

Numerous investigators have attempted to identify the physiological functions of epicardial fat (Figure 1) [9]. Firstly, EAT is a vital energy lipid storage for myocardium in periods of elevated energy requirements [21]. It has been demonstrated that epicardial fat uptakes and releases free fatty acids (FFA) to a much greater degree than other visceral adipose tissues [22]. Free fatty acids are the dominant energy source for a healthy heart’s contraction, whereas their oxidation covers up to 70% of the heart’s total energy requirements [22]. Impressively, fatty acid-binding protein 4 (FABP4) is extensively expressed by human epicardial adipocytes [23]. This protein transfers free fatty acids from EAT to the adjacent heart muscle via either vasocrine or paracrine routes [23]. Moreover, epicardial fat quickly utilizes redundant free fatty acids since it converts them into lipid storage units for future energy myocardial demands. [14]. Thus, myocardium may be efficiently protected from lipotoxicity [11].
FIGURE 1: Physiological functions of epicardial adipose tissue (EAT)

EAT: Epicardial adipose tissue, FFA: Free fatty acid, BAT: Brown adipose tissue

This figure was created by the author of the article, Nikoleta Karampetsou.

Mechanical Protection

Due to its location in the atrioventricular and interventricular chambers and its vicinity to coronary vessels, EAT serves as a cushion [24]. In this way, it protects coronary arteries from violent distortion triggered by arterial pulse wave and myocardial contraction [24]. Furthermore, it acts as a supplemental anatomic layer for the total heart mass in case of mechanical injury [24].

Brown Adipose Tissue Properties

Brown adipose tissue (BAT) is a subject of vigorous scientific investigation since its role in human beings has not yet been clarified [25]. Protection against low temperatures via thermoregulatory mechanisms and stressful hemodynamic states, such as ischemia and hypoxia are the main functions of brown fat [25]. It has been described that in neonates epicardial brown fat cells are the majority of EAT adipocytes [26]. However, from infant to adult life the number of brown adipocytes annually decreases [26]. Recently, it has been proposed that EAT could be defined as beige adipose tissue, since brown fat cells are gradually replaced by small unilocular white adipocytes, similar to beige lineage ones [25]. Uncoupling protein-1 (UCP-1) and other brown specific proteins are impressively greatly expressed in human EAT [25]. Interestingly, UCP-1 presents higher concentration in human EAT than in other visceral fat units [27]. Brown specific fat proteins facilitate heart thermoregulation and protect it against stressful conditions, as mentioned above [25]. According to Sacks et al., advanced coronary atherosclerosis is associated with the elimination of these proteins in favor of significant elevation of inflammatory ones [28].

Epicardial Adipose Tissue as an Endocrine Organ

Epicardial adipose tissue is not only an energy fat source but also an active endocrine organ with several vasocrine and paracrine effects [29]. Under physiological circumstances, epicardial fat secretes plenty of beneficial cytokines, such as adiponectin, adrenomedullin and omentin-1 which present anti-atherogenic, anti-inflammatory and antithrombotic properties [29]. Paracrine and vasocrine pathways have been suggested as the two main interaction routes between EAT and the human heart [15]. The paracrine method promotes the adipokines’ diffusion from the pericoronary adipose tissue to the arterial wall and the direct interaction with the endothelial and vascular smooth cells [30]. The vasocrine way includes the direct transit of free fatty acids and adipokines into the coronary lumen’s vasa vasorum [31]. The adipocytokines mentioned above activate adenosine monophosphate-activated protein kinase (AMPK), an enzyme mostly produced by heart and skeletal muscles [32, 33]. Under cellular stress states and low energy supplies, AMPK induces catabolic pathways and restricts anabolic ones in order to increase cellular adenosine triphosphate (ATP) levels and regulates glucose and free fatty acid uptake [32,33].

Adiponectin is a protein hormone mainly produced by adipocytes. Epicardial fat expresses adiponectin receptor 1 (AdipoR1) and adiponectin receptor 2 (AdipoR2) [34]. Adiponectin connection to these receptors leads to free fatty oxidation, triacylglycerol (TAG) turnover, and a decrease in fat storage in human EAT [35]. Secondly, through AMPK activation adiponectin restricts the production of acetyl coenzyme A (acetyl-CoA) [35]. In this way, both glucose levels and fatty oxidation are regulated [35]. Noticeably, adiponectin also induces an anti-inflammatory microenvironment in the human heart since it inhibits the production of...
inflammatory cytokines, such as tumor necrosis factor-alpha (TNF-α) and interleukin-6 (IL-6) [36].

Adrenomedullin is a protein which presents crucial cardioprotective features. It interacts with the calcitonin receptor-like receptor (CRLR) and stimulates protein kinase A (PKA) in cardiac muscle cells [37]. Therefore, calcium channels are activated and calcium is released in cardiomyocytes cytoplasm leading to the elevation of cardiac output [38]. Except for this function, it has been reported that adrenomedullin inhibits endothelial cell apoptosis and reduces oxidative stress [38]. Another adipocytokine that is highly expressed in EAT under normal circumstances is omentin-1 [36]. It is characterized by anti-atherogenic properties since it induces the activation of anti-inflammatory M2 macrophages and subdues foam cell formation linked to oxidization of low-density lipoprotein (LDL) [39]. In their study, Du et al. demonstrated that omentin-1 expressed in EAT was significantly reduced in patients with CAD than in individuals without [36]. Finally, it is now evident that EAT in comparison to subcutaneous adipose tissue (SAT) releases greater levels of proteins associated to potassium channel interaction, biosynthesis and wound healing [40].

Epicardial adipose tissue: Assessment techniques

Nowadays, growing evidence suggests that excessive epicardial adiposity is an independent risk factor for cardiovascular disease development [41]. Increased EAT thickness and volume are clinical markers of excessive visceral fat accumulation [41]. Thus, accurate quantification of EAT could serve both as a crucial prognostic tool and as a medication target for cardiovascular diseases, especially CAD [41]. In 2003, Iacobellis et al. first assessed EAT thickness through two-dimensional echocardiography [42]. Epicardial fat appears as an echo-free space above the right ventricle’s free wall, where it is more prominent [42]. Parasternal long-axis ultrasound images in combination with short-axis ones at the end of systole during three cardiac cycles are performed in order to achieve precise measurement of EAT [42]. To our knowledge, 5 mm of epicardial fat thickness recorded in transthoracic echocardiography has been proposed as a cut-off value and prognostic risk factor for CAD [10]. Transthoracic echocardiography remains an easy, economic, noninvasive and easily reproduced procedure of EAT quantification [11]. However, total eat volume cannot be estimated by 2D-echo [11]. Additionally, the quality of measurements highly depends on the operator’s experience [11, 42].

Cardiac magnetic resonance imaging (MRI) and multislice computed tomography (MCT) are novel and promising imaging techniques for reliable volumetric assessment of epicardial fat [31]. The MRI provides explicit information regarding total EAT volume and mass by using the spin-echo sequence technique and without radiation exposure to patients [43]. Specifically, manual contouring of epicardial fat region in short-axis views at the end of diastole during several cardiac cycles provides precise quantification of EAT volume in each slice [31]. The total of the above measurements indicates the total EAT volume in the human heart [31]. Despite the advantages, MRI is still an expensive and hardly accessible method in daily clinical practice [44].

On the other hand, either contrast or non-contrast enhanced MCT is the most desirable EAT imaging method, since it has better spatial resolution and not only identifies exact total EAT volume but also evaluates coronary arterial calcification [31,45]. In addition to this, cardiac computed tomography angiography (CCTA) is a similar method which not only estimates coronary artery stenosis but also recognizes dangerous atherosclerotic plaque characteristics, such as vulnerable atheromas [46]. Consequently, it may be suggested as a valuable risk stratification tool [46]. However, the cost, the restricted availability and the remarkable exposure to ionizing radiation are major disadvantages of both MCT and CCTA [45, 47].

Despite its several limitations, two-dimensional echocardiography remains the most preferable and cost-effective method of epicardial fat evaluation in daily clinical practice (Table 1).
Epicardial adipose tissue and coronary artery disease

Excessive adiposity has a well-established role in coronary atherosclerosis by both secreting plenty of active cytokines and regulating insulin sensitivity [48]. During the last two decades, interest regarding EAT and CAD has been growing rapidly [12]. Unobstructed anatomical proximity of EAT and coronary arterial tree indicates the substantial role of it in coronary atherosclerosis progression [8]. In addition to local vicinity, however, EAT seems to induce CAD in numerous ways [8].

Coronary artery disease is inflammatory in nature [3]. Inflammatory properties of epicardial fat have been investigated since the early 2000s [11]. Multiple studies have reported dense infiltration of inflammatory cells in human EAT [11]. In 2011, Hirata et al. demonstrated a greater concentration of inflammatory M1 macrophages than anti-inflammatory M2 macrophages in EAT samples derived from patients suffering from advanced CAD [49]. It is now well-known that the increased prevalence of M1 macrophages persuades coronary atherosmas instability and thereafter rupture [49]. Except for macrophages, the presence of mast cells, B lymphocytes, T lymphocytes, and dendritic cells in EAT has also been associated with coronary atherosclerosis [50]. All the previously mentioned cells express in their membranes Toll-like receptors (TLRs) [51]. Extracellular ligands, such as saturated fatty acids connect to TLRs and promote immune response [51]. Particularly, nuclear factor-κB (NF-κB) and JUNN-terminal kinase (JNK) are activated and induce the upregulation of inflammatory molecules in EAT [51]. In their study, Baker et al. demonstrated elevated activation of NF-κB and JNK pathophysiology pathways in EAT biopsies of people suffering from advanced CAD [51].

In a benchmark study, Mazurek et al. observed elevated concentrations of inflammatory mediators in human EAT [15]. Interleukin-6 (IL-6), interleukin-1 (IL-1), and TNF-α were all upregulated in subjects suffering from CAD [49]. In addition to this, the inflammation level was higher in epicardial fat than subcutaneous adipose tissue in these subjects [15]. High TNF-α concentrations increase IL-6 levels [52]. Both of them elevate endothelial and vascular smooth cell permeability, while they also have a negative influence on insulin tissue sensitivity [52]. Since then, several studies have identified genes encoding other inflammatory adipokines, such as resistin and chemerin [56,52]. Due to the lack of anatomic intervening barrier, inflammatory cytokines expressed by EAT affect coronary arteries through paracrine and vasocrine pathways [53]. Lately, it has been shown that elevated abnormal EAT proteasome leads to thicker EAT layer, more intense chronic inflammation and therefore more severe CAD [15,44,54]. By contrast, cardioprotective cytokines, such as adiponectin and adrenomedullin were found extremely low in EAT samples from people with CAD [11]. The reduction of such anti-inflammatory molecules leads to further coronary inflammation [11]. Finally, leptin is another highly expressed hormone in the pericoronary EAT [55]. Leptin expression is upregulated by IL-6 and downregulated by TNF-α [56]. Increased leptin levels promote endothelial cell permeability and induce monocytes and macrophages adhesion, and destabilize atheromatous plaques in the coronary lumen [55]. It has been noticed that the adiponectin/leptin ratio in EAT decreases under pathological conditions and is a novel risk factor for CAD [55,57].

Epicardial adipose tissue has also been gaining interest regarding oxidative stress [53]. Recently, it has been proposed that epicardial fat in humans suffering from CAD presents higher concentration of reactive oxygen species (ROS) [58]. In their study, Demir et al. found increased total oxidative stress (TOS) in epicardial fat compartment of patients suffering from CAD, and metabolic syndrome [59]. The ROS activates various transcription factors which result in messenger ribonucleic acid (mRNA) encoding of genes involved in local inflammation [59]. Cellular stress mediators, such as mitogen-activated protein kinase kinase kinase 5 (MAP3K5) are highly expressed by epicardial adipocytes leading to cellular apoptosis and further endothelial dysfunction [40]. In this way, the imbalance between ROS and inflammatory factors results in chronic

| Imaging techniques       | Availability | Cost | Radiation | Epicardial adipose tissue (EAT) thickness assessment | EAT volume assessment | Coronary artery calcification |
|--------------------------|--------------|------|-----------|-----------------------------------------------------|-----------------------|-------------------------------|
| Echocardiography         | easily available | low  | no        | yes                                                 | no                    | no                            |
| Magnetic resonance imaging (MRI) | not easily available | very high | no | yes                                                 | yes                   | no                            |
| Computed tomography (CT) | not easily available | high | yes       | yes                                                 | yes                   | yes                           |

TABLE 1: Pros and cons of main current imaging methods for epicardial adipose tissue (EAT) quantification

EAT: Epicardial adipose tissue, MRI: Magnetic resonance imaging, CT: Computed tomography
inflammation, endothelial dysfunction, and consequently, coronary atherosclerosis. [40]

Lipotoxicity is deemed to be another pathophysiology mechanism which promotes EAT-induced coronary atherosclerosis [60]. Nowadays, it is known that excessive epicardial fat provokes imbalance in lipid and glucose metabolism [60]. Redundant epicardial fat secretes a high amount of free fatty acids, which are then accumulated in coronary artery lumen promoting atheromatous plaque development [10,13]. Remarkably, phospholipase A2 and endothelial lipase are significantly elevated in EAT of individuals with CAD [66]. These proteins participate in lipid metabolism pathways and could serve in further ectopic fat accumulation in adjacent coronary vessels [60]. Furthermore, epicardial fat is deemed to be an insulin-resistant lipid compartment [61]. In fact, it is characterized by lower glucose utilization rate than other visceral fat tissues [61]. Glucose-transporter 4 (GLUT4) is an intracellular protein mainly expressed in muscle cells and adipocytes [62]. It regulates glucose uptake in fat and muscle cells after insulin cascade activation [62]. Epicardial adipocytes are generally characterized by lower levels of glucose transporter type 4 (GLUT-4) [32]. However, in diabetic individuals also suffering from coronary atherosclerosis, GLUT4 levels in EAT were even lower, underlining in this way the local role of insulin-induced atherogenesis in CAD [62].

All these pathophysiology pathways underlie a direct relationship between epicardial fat and CAD [9]. Consequently, EAT may be used as a screening and risk stratification tool [48]. Both EAT thickness and volume have been measured higher in humans with CAD than in those without [13]. In their clinical investigation, Fahri et al. correlated elevated mean EAT thickness with critical atheromatosis as demonstrated by Syntax and Gensini scores [63]. Increased coronary artery calcium score (CACS >10) was also observed in patients with high EAT volume as measured by either CT or MRI [64]. Recently, Cosson et al. proposed the quantification of EAT volume as an independent parameter in the CACS determination in diabetic patients [65]. In another study, elevated EAT volume was tightly associated with vulnerable atherotic plaques in patients suffering from symptomatic atherosclerosis but had zero CACS [66]. Taking this into consideration, the EAT assessment may be used for early diagnosis of non-calcified and unstable atheromatous plaques [66]. Moreover, epicardial fat is not equally distributed in the whole heart mass [57]. Because of their anatomic proximity, coronary arteries are mainly affected by pericoronary EAT [64]. A higher pericoronary EAT volume has been referred as a determining risk factor leading to more severe CAD and calcification in postmenopausal women [67]. Same results were also observed in prediabetic patients with history of myocardial infarction [57]. Lastly, the increased levels of inflammatory cytokines in serum and EAT samples of humans with CAD underlie the crucial role of abnormal epicardial fat proteasome in coronary atherosclerosis progression. [18,29,30]

**Therapeutic targets in epicardial adipose tissue**

*Lifestyle Modifications*

Undoubtedly, epicardial fat is a novel cardiovascular risk factor [29]. Despite the vigorous investigation of the main pathogenic mechanisms of EAT, there is still little knowledge regarding suitable medical treatment of this pathological epicardial fat compartment (Figure 2) [29]. Lifestyle modifications, such as weight loss, aerobic and anaerobic physical exercise seem to be key factors in achieving EAT thickness reduction [11]. Iacobellis et al. noticed a significant decrease in EAT thickness of critically obese patients (BMI 45± 5 kg/m²) who attended a low-calorie diet for up to six months [68]. In a recent pilot study, Konwerski et al. investigated the effect of intense aerobic physical exercise on epicardial fat volume and metabolic profile among 30 ultramarathon amateur athletes and eight individuals with a sedentary lifestyle (control group) [69]. The EAT volume as measured by cardiac MRI was significantly lower in ultramarathon athletes than in the control group [69]. Furthermore, ultra-runners also presented better metabolic profile (lower lipid concentration and lower BMI) as well as lower levels of plasma IL-6 when compared to the control group [69]. In a recent meta-analysis of 10 studies, similar positive effects of endurance training on EAT were also noticed [70]. Additionally, it is widely known that obesity is associated with coronary atherosclerosis and excessive visceral adiposity, including increase in total EAT mass [71]. In 2018, Altin et al. investigated the potential role of bariatric surgery, and specifically sleeve gastrectomy, in epicardial fat thickness. They reported the eliminated EAT thickness in all echocardiography measurements after bariatric surgery [71].
Medication Treatment

With the exception of lifestyle modifications, several medications have been under investigation in order to indicate their probable role in abnormal EAT alteration [72]. Antilipidemic drugs such as statins not only reduce EAT thickness but also restrict pathological metabolic activity of epicardial fat due to their pleiotropic properties [72]. Firstly, statins inhibit competitively the transformation of hydroxymethyl glutaryl-CoA to mevalonic acid, resulting in increased expression of LDL receptors. Furthermore, statins present anti-inflammatory features since they restrict the growth of macrophages, reduce immune cell adhesion in atheromatous plaques and suppress proinflammatory cytokines secretion [73]. Raggi et al. formed a randomized controlled trial for one year with 420 postmenopausal women, in which 194 were treated with 80 mg atorvastatin whereas the other 226 received 40 mg pravastatin. All the patients underwent chest computed tomography in order to quantify epicardial fat attenuation in Hounsfield units (EAT HUs). The EAT HUs were measured in the area of right coronary artery. Atorvastatin reduced EAT volume more efficiently than pravastatin. However, both of them decreased EAT HUs almost similarly, demonstrating the anti-inflammatory properties of statins [74]. Proprotein convertase subtilisin/kexin type 9 (PSCK9) is an inflammatory mediator highly produced by visceral fat depots, especially EAT [75]. It promotes arterial atherosclerosis through direct interaction with LDL receptors and monocytes adhesion in the atheromatous plaque [75]. The PCSK9 inhibitors are lipid-lowering drugs which may be involved in the reduction of EAT thickness [76]. In 2020, Galvez et al. observed that in 24 patients who received either evolocumab or alirocumab for six months, EAT thickness was significantly reduced as measured by transthoracic echocardiography [76].

To the best of our knowledge, several medications used for diabetes and/or obesity may also provide cardioprotective benefits beyond glycaemic control [77]. Metformin remains the most commonly administered hypoglycemic substance worldwide [78]. It is proposed as the gold-standard for initial therapy of type-2 diabetes [78]. Several studies have demonstrated its determining role in blood glucose reduction and increase in insulin sensitivity in target organs, such as visceral fat [57]. In 2019, Ziyrek et al. reported that the initiation of metformin in 40 diabetic subjects resulted in significant decrease in EAT thickness three months after [78]. Pioglitazone belongs to a family of hypoglycemic drugs called thiazolidinediones [79]. It stimulates both peroxisome proliferator-activated receptor gamma (PPAR-γ) and proliferator-activated receptor alpha (PPAR-α) [79]. Thus, it inhibits the secretion of pro-inflammatory cytokines in visceral fat depots, alternating the metabolic profile [28]. Grosso et al. reported important depletion of cytokines (IL-6, TNF-α, resistin, etc.) expressed in EAT biopsies derived from patients who underwent cardiovascular bypass graft surgery (CABG) and were treated with pioglitazone or pioglitazone/simvastatin compared to the control group [79]. Dipeptidyl peptidase 4 (DPP-4) inhibitors are oral antidiabetic medications which regulate blood glucose levels by stimulating incretin production, inhibiting glucagon release and increasing insulin levels [80]. Sotagliflozin is a DPP-4 inhibitor which has been investigated regarding its role in abnormal epicardial fat modulation [80]. Lima-Martínez et al. conducted an
interventional study on 26 overweight and diabetic patients insufficiently treated with metformin [80]. The combination of sitagliptin/metformin at a dosage 50/1000 twice a day was registered to these subjects [80]. Epicardial fat was assessed before and after this medical intervention via ultrasound [80]. Significant reduction (from 9.98 ± 2.63 to 8.10 ± 2.11 mm, p = 0.001) in EAT was achieved after sitagliptin admission [80]. Glucagon-like peptide-1 (GLP-1) receptor agonists are injectable drugs which not only attenuate diabetic hyperglycaemia but also reduce obesity and major adverse cardiovascular events [81]. Epicardial adipocytes express GLP-1 receptor that has been shown to inhibit local adipogenesis by increasing free fatty acid oxidation to induce brown adipose tissue differentiation and to increase insulin sensitivity [75,82].

Current research has shown positive results in EAT thickness-reduction in diabetic patients who received either liraglutide, semaglutide, or dulaglutide [83,84]. Sodium-glucose co-transporter -2 (SGLT-2) inhibitors are novel hypoglycemic agents which block glucose reabsorption in the proximal kidney tubule and consequently result in glucosuria and serum glycemic control [85]. Administration of such agents leads to weight loss and decline in visceral adiposity [83]. Particularly, SGLT-2 inhibitors-induced hypoglycemia promotes lipolysis, free fatty acid oxidation and improved visceral lipid and glucose metabolism [86]. Noticeable reduction of EAT volume, serum inflammatory cytokines and increase in adiponectin levels were identified in 40 diabetic individuals with CAD after the administration of dapagliflozin for six months, indicating a positive role of SGLT-2 in EAT function and overall metabolic profile [87]. Similar results were also noticed after the administration of empagliflozin [88]. Insulin restoring treatment is also associated to EAT thickness reduction [89]. In 2016, Elisha et al. denoted a clear decline in epicardial fat composition in diabetes mellitus 2 patients who were treated with either detemir or glargine [89]. However, the insulin effect was more prominent in the detemir group [89].

Conclusions
Nowadays, special attention is paid to the complex physiology and pathophysiology properties of epicardial fat. Epicardial adipose tissue is now considered a determining cardiometabolic risk factor for cardiovascular diseases, especially CAD. Multiple studies have been conducted to investigate the plausible pathophysiology pathways regarding abnormal EAT and coronary atherosclerosis progression. Elevated secretion of inflammatory molecules and significant reduction of anti-inflammatory ones, the elevation of immunity cells reaction, increased levels of reactive oxygen species, lipotoxicity and glucotoxicity have been proposed as the main etiopathogenic routes. Taking all these pleiotropic effects of epicardial fat into account, accurate measurement of this fat unit seems to be essential. Imaging techniques, such as conventional echocardiography, CT, and MRI should be easily accessible in everyday clinical practice for the measurement of EAT. To the best of our knowledge, both lifestyle and pharmacological interventions could modulate abnormal EAT volume and thickness, restore its cardioprotective effects and eliminate coronary atherosclerosis progression. A healthy lifestyle and medical treatment with either traditional or novel antidiabetic and lipid-lowering agents have shown encouraging results. More clinical investigations should be conducted to better understand the exact pathophysiology mechanisms of EAT and determine whether the pharmacological alteration of its mass could efficiently prevent CAD.

Additional Information
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