Review Article

Update on RAAS Modulation for the Treatment of Diabetic Cardiovascular Disease

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Since the advent of insulin, the improvements in diabetes detection and the therapies to treat hyperglycemia have reduced the mortality of acute metabolic emergencies, such that today chronic complications are the major cause of morbidity and mortality among diabetic patients. More than half of the mortality that is seen in the diabetic population can be ascribed to cardiovascular disease (CVD), which includes not only myocardial infarction due to premature atherosclerosis but also diabetic cardiomyopathy. The importance of renin-angiotensin-aldosterone system (RAAS) antagonism in the prevention of diabetic CVD has demonstrated the key role that the RAAS plays in diabetic CVD onset and development. Today, ACE inhibitors and angiotensin II receptor blockers represent the first line therapy for primary and secondary CVD prevention in patients with diabetes. Recent research has uncovered new dimensions of the RAAS and, therefore, new potential therapeutic targets against diabetic CVD. Here we describe the timeline of paradigm shifts in RAAS understanding, how diabetes modifies the RAAS, and what new parts of the RAAS pathway could be targeted in order to achieve RAAS modulation against diabetic CVD.

1. Introduction

Cardiovascular diseases (CVD) are the main cause of diabetes-related morbidity and mortality [1, 2]. They include myocardial infarction, which is due to premature atherosclerosis, and diabetic cardiomyopathy, both leading to heart failure. Patients with diabetes have a higher prevalence of cardiovascular morbidity and mortality as compared to the general population [3], such that diabetes is considered not only as an independent cardiovascular risk factor but also as a cardiovascular event equivalent, meaning that patients with diabetes have a risk of cardiovascular complications equal to that of patients with a prior myocardial infarction [4]. This excess cardiovascular risk in comparison to the general population is explained only partly by conventional cardiovascular risk factors, such as hyperglycemia, dyslipidemia, hypertension, and cigarette smoking.

One of the links between diabetes and such a high prevalence of CVD is renin-angiotensin-aldosterone system (RAAS) activation. It has been shown that the RAAS plays a major role in the development of diabetic cardiovascular complications [5], as it promotes atherosclerosis [6, 7], cardiomyocyte loss, and extensive myocardial fibrosis [8, 9]. Consistent with this view, ACE inhibitors and angiotensin II receptor blockers represent the first line therapy for primary and secondary CVD prevention in patients with diabetes [10]. Recent research has uncovered new dimensions of the RAAS and, therefore, new potential therapeutic targets against diabetic CVD. Here we describe the timeline of paradigm shifts in RAAS understanding, how diabetes modifies the RAAS, and what new parts of the RAAS pathway could be targeted in order to achieve RAAS modulation against diabetic CVD.

2. Paradigm Shifts in the Renin-Angiotensin-Aldosterone System Understanding

2.1. The Renin-Angiotensin-Aldosterone System Has Hemodynamic and Nonhemodynamic Actions. The renin-angiotensin-aldosterone system (RAAS) consists of a group
of enzymes and peptides whose main function is to control blood pressure by regulating vasoconstriction, sodium reabsorption, and body fluid homeostasis. The modern view of the RAAS began with the concept that this was a life-saving system, which raised blood pressure by approximately 30 mmHg in case of an acute hemorrhage [11]. Classically, the process whereby the RAAS raises blood pressure usually starts within the kidney, where a blood pressure fall stimulates renin release into the bloodstream [12, 13]. Then, circulating renin cleaves hepatic angiotensinogen and generates angiotensin (Ang) I, which is converted to Ang II by pulmonary angiotensin-converting enzyme (ACE), as represented in Figure 1 [14–16]. Soon after its generation, Ang II causes smooth muscle cell vasoconstriction, stimulates the sympathetic nervous system, and promotes renal retention of salt and water by binding to its specific receptors [17, 18]. Moreover, in the adrenal glands, Ang II stimulates the release of aldosterone, which enhances tubular sodium reabsorption in the kidney and increases the effective circulating plasma volume [19].

Ang II has two main receptors: Ang II type 1 receptor (AT1R) and Ang II type 2 receptor (AT2R), as represented in Figure 2. Studies in mice lacking AT1R showed that the hemodynamic actions of Ang II depended on AT1R [20–22]. On the other hand, AT2R, which was found highly expressed in differentiated mesenchymes during fetal life and decreased rapidly after birth, seemed to regulate fetal development [18, 23]. Nevertheless, AT2R is still detectable in adulthood in different organs, such as the heart, kidney, adrenal glands, brain, ovaries and uterus, and the vessels [24], where its main function is to counterbalance AT1R. For instance, AT2R-knockout mice exhibit high blood pressure levels and increased vascular sensitivity to Ang II [25, 26], suggesting that AT2R antagonizes AngII-AT1R peripheral effects. Such an action could be achieved not only by vasodilator effects that are independent of AT1R [27, 28] but also by AT1R downregulation [29] and/or direct inhibition of AT1R signaling [30, 31], as represented in Figure 2.

Although the RAAS was a life-saving system, the first aspect that became clear was that if the Ang II cascade was activated inappropriately, it could lead to hypertension and CVD [32]. Therefore, ACE inhibitors and Ang II type 1 receptor blockers (ARB) were designed as antihypertensive therapies [33, 34]. A paradigm shift in RAAS understanding occurred when the CONSENSUS and the SOLVD trials showed that the ACE inhibitor Enalapril reduced overall mortality by 27% and 16% in patients with heart failure [35, 36]. Enalapril’s benefit suggested that the RAAS had significant nonhemodynamic actions that were associated with cardiovascular pathology. These trials led to mechanistic studies demonstrating that Ang II promoted ventricular hypertrophy [37–39], myocardial infarction [40], and atherosclerosis [41–43], independent of blood pressure values. In particular, by binding to AT1R, Ang II was able to induce reactive oxygen species generation, tissue inflammation and fibrosis, and the regulation of cell growth and differentiation as well as apoptosis and survival as summarized in Table 1 [41–63].

2.2. Circulating and Tissue Renin-Angiotensin-Aldosterone System. The observation that many tissues were capable of synthesizing the RAAS key components led to another paradigm shift in the timeline of RAAS understanding: the RAAS was not anymore only a circulating hormonal system but also a tissue system widespread in cardiovascular organs [64–68]. ACE and Ang II receptors were identified in all the relevant target tissues including the heart, kidney, blood vessels, and adrenal glands [69–71], where they have not only endocrine but also paracrine and autocrine effects [72]. Interestingly, circulating and local systems can sometimes behave in opposite ways, like in cases of high salt intake, which generally downregulates circulating and upregulates local RAAS [73].

Renin, angiotensinogen, ACE, and Ang II receptors were all present in the heart [67], where they were found upregulated in models of cardiac injury, such as volume overload [74], myocardial infarction [75], and heart failure [76–78]. For example, one of the first experimental observations was that ACE increased in heart failure, its cardiac induction was
Angiotensin II has two major receptor isoforms: Ang II type 1 receptor (AT1R) and Ang II type 2 receptor (AT2R). AT1R stimulation mediates the classical actions of Ang II, including hemodynamic and nonhemodynamic effects, leading to hypertension, cardiac remodeling, and atherosclerosis. On the other hand, AT2R stimulation usually causes opposing effects to AT1R. Moreover, it can antagonize AT1R by downregulating it, inhibiting its signaling, or binding to it. Ang is for angiotensin; AT1R is for Ang II type 1 receptor; AT2R is for Ang II type 2 receptor; NO is nitric oxide.

Almost a decade after the CONSENSUS and the SOLVD studies, the HOPE [5] trial extended to high-risk patients, such as those with diabetes, the concept that RAAS blockade protected against cardiovascular morbidity and mortality. In that trial, Ramipril significantly reduced the rates of death, myocardial infarction, and stroke in patients with vascular disease or diabetes. Likewise, the LIFE trial showed that Losartan was more effective than Atenolol in reducing cardiovascular morbidity and mortality in patients with hypertension, diabetes, and ventricular hypertrophy [80].

At the same time the RALES and EPHESUS trials showed that aldosterone antagonism on top of Ang II blockade provided a major additive benefit as it reduced significantly overall mortality and the rate of death from cardiovascular causes among patients with severe heart failure [84] or after acute myocardial infarction [85]. These results suggested that Ang II blockade could not totally suppress the production of aldosterone and that other factors in addition to Ang II were important in the production of this hormone. Moreover,
they reinforced the notion that aldosterone promoted organ damage by a series of cellular and tissue effects, elicited by binding predominantly to the mineralocorticoid receptors, which are reported in Table 1 [86–103]. More recently, Eplerenone benefits have been extended to diabetic patients, where this treatment reduced adverse CV events in diabetic patients with heart failure following myocardial infarction [104]. A randomized study evaluating the effects of two aldosterone antagonists, Finerenone versus Eplerenone, in patients with heart failure following myocardial infarction [104]. A randomized study evaluating the effects of two aldosterone antagonists, Finerenone versus Eplerenone, in diabetic patients with heart failure is still ongoing [105].

2.3. ACE2 and (Pro)Renin Discovery. Nevertheless, ACE2 discovery represents perhaps the last fundamental paradigm shift in RAAS understanding [106, 107]. ACE2 is a carboxypeptidase whose main function is to degrade Ang II to generate Ang 1–7 [108, 109]. Ang 1–7 is a biologically active peptide, which exerts opposite peripheral actions to those of Ang II [110] by binding predominantly to the Mas1 receptor (Mas1R) [111]. Cellular and tissue effects of Ang 1–7/Mas1R are summarized in Table 1 [112–128]. Moreover, Ang 1–7, like all Ang II "breakdown" products, has the potential to act as an endogenous ligand of AT2R [129], further promoting organ protection (Figure 3).

The identification of ACE2 provided evidence that the RAAS had two pathways with opposite effects: the classic ACE/Ang II/AT1R and the new ACE2/Ang 1–7/Mas1R (and AT2R) pathway. Accordingly, the current scientific opinion is that what is critical in CVD development is an imbalance in ACE-Ang II and ACE2-Ang 1–7 [130]. Consistent with this view, ACE2 is regarded as the central regulator of the RAAS [131], as its changes can affect not only Ang II detrimental actions but also Ang 1–7 protective effects. For instance, a decrease in ACE2 results in activation of the Ang II/AT1R axis with a parallel reduction of Ang 1–7 [51, 132]. Nevertheless, ACE2 protective effects can be attributed not only to the degradation of Ang II and the generation of Ang 1–7 but also to the degradation of Ang I and the generation of Ang 1–9 [106], as represented in Figure 3, but also to the local increase of ANP. Recently, we have shown that ACE2 regulates ANP production through Ang 1–7 [132].

Having said that, the final entry in the RAAS is the (pro)renin receptor [(P)RR], which is a specific receptor for both renin and its inactive precursor prorenin [133], collectively called (pro)renin (Figure 3). When (pro)renin binds to (P)RR it can degrade angiotensinogen to Ang I, but it can also trigger intracellular signaling pathways, which are independent of Ang II generation [133]. Interestingly, the ability of prorenin to activate the RAAS depends on the fact that its binding induces a conformational change that exposes

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<th>Table 1: Cellular and tissue effects of Ang II, Ang 1–7, aldosterone, and (pro)renin in normal conditions.</th>
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* Cellular and tissue effects are predominantly mediated by the indicated receptors.
Ang is for angiotensin; AT1R is for Ang II type 1 receptor; Mas1R is for Mas1 receptor; MR is for mineralocorticoid receptor; and (P)RR is for (pro)renin receptor.
Figure 3: Tissue renin-angiotensin-aldosterone system. The activation of the RAAS cascade begins with renin secretion. Renin is secreted as a precursor protein, prorenin. Renin and prorenin can bind to their specific receptor called (P)RR and activate intracellular pathways independent of RAAS. Otherwise, renin cleaves angiotensinogen to form Ang I, which is then converted to Ang II by ACE. Ang II binds to its specific receptors: AT1R and AT2R. AT1R promotes blood pressure increase, cardiac remodeling, and atherosclerosis development. In addition, through AT1R, Ang II stimulates aldosterone secretion. On the other hand, AT2R seems to antagonize these effects. Moreover, Ang I and Ang II can be cleaved by ACE2 to form Ang 1–9 and Ang 1–7, which have opposite effects to those of Ang II, such as vasodilation, anti-inflammatory, antifibrotic, and antiremodeling effects, which are mediated by Mas1R and partly by AT2R. In addition, Ang 1–7 stimulates local ANP secretion. ACE is for angiotensin-converting enzyme; ACE2 is for angiotensin-converting enzyme 2; Ang is for angiotensin; AT1R is for AngII type 1 receptor; AT2R is for Ang II type 2 receptor; ANP is for atrial natriuretic peptide; (P)RR is for (pro)renin receptor.

prorenin active site, which becomes catalytically active and generates Ang I. The biology of (P)RR is very complex and only partially understood [134]. The association between (P)RR gene polymorphism and ambulatory BP in Japanese men [135] has suggested that (P)RR was related to cardiovascular disease leading to experimental studies supporting this hypothesis. The cellular and tissue effects of (pro)renin are reported in Table 1 [136–145]. Nevertheless, transgenic animals overexpressing (pro)renin developed hypertension that was sensitive to ACE inhibition, indicating that this effect was due to Ang II generation, which led to arguing against an independent role of (P)RR in CVD [146, 147]. On the other hand, Batenburg and Danser have suggested that (pro)renin overexpression led to a 400-fold increase of plasma (pro)renin levels, which is far from the 1000-fold–100 000-fold increase that is required to observe (pro)renin effects in vitro [148]. Certainly, (P)RR has essential functions that are independent of the RAAS and are related to the vacuolar H⁺-proton adenosine triphosphatase (V-ATPase) [149]. In particular, (P)RR is a cofactor of the V-ATPase [150] and is required for central nervous system development [151, 152]. (P)RR is also an essential partner in Wnt receptor complex signaling [153], which regulates normal patterning of the embryo, and in adults, cell proliferation, migration, polarity, and tissue repair. These findings explain not only why (P)RR deletion yielded embryos that died before the end of embryogenesis [153, 154] but also why mice with specific ATP6ap2/(P)RR knockout in cardiomyocytes died of severe heart failure within 3 weeks of age [155].

3. Diabetes, the Activation and Imbalance of the Renin-Angiotensin-Aldosterone System, and Cardiovascular Disease

3.1. The Renin-Angiotensin-Aldosterone System Activation Links Diabetes to CVD. The current picture of the RAAS is that of an extremely complex pathway whose overstimulation triggers a chain of events contributing to cardiovascular disease. The RAAS is found not only in the circulation, where it has hemodynamic effects, but also at a tissue level, where it has nonhemodynamic effects. It is activated in the hypertrophied, ischemic, and failed heart, and activation of the ACE/AngII/AT1R axis leads to both accelerated atherosclerosis and direct cardiac injury, as shown by infusing Ang II or by the studies where RAAS blockade was able to attenuate or prevent cardiac damage [78].

Patients with diabetes have a higher prevalence of cardiovascular morbidity and mortality as compared to the general
population [3]. On one hand, diabetes is associated with accelerated atherosclerosis affecting the coronaries, which increases the risk for myocardial infarction and heart failure. On the other hand, it can cause diabetic cardiomyopathy, which corresponds to a cardiac dysfunction independent of coronary artery disease, hypertension, and valvular complications [156].

Upregulation of the RAAS is evident in diabetes. Miller and colleagues demonstrated that during the early stages of diabetes there was an increase in plasma renin activity, mean arterial pressure, and renal vascular resistance [157]. Moreover, the fact that Losartan lowered blood pressure more in hyperglycemic than in euglycemic conditions [158], and that Captopril and Eprosartan caused a greater renal vasodilator response during hyperglycemia [159], suggested that glucose levels were associated with an activation of the RAAS, which made diabetic patients more sensitive to RAAS antagonism. Consistent with this view, direct renin inhibition with Aliskiren led to significant improvement of left ventricular hypertrophy [160] and end-systolic volume [161] only in groups of patients with diabetes.

The importance of RAAS antagonism in the prevention/reduction of cardiovascular complications has clearly demonstrated that diabetes-induced RAAS activation contributes substantially to diabetic CVD [5, 80–82]. Activation of AngII/AT1R pathway in the setting of diabetes can in fact promote cell growth and proliferation, apoptosis [162], oxidative stress generation [9], inflammation [6], and fibrosis [9], which are all leading to cardiac remodeling and atherosclerosis, that can be reversed/reduced by RAAS blockade [7–9].

3.2. Diabetes-Induced Activation of the AngII/AT1R Pathway and Imbalance of the Renin-Angiotensin-Aldosterone System

There are several mechanisms whereby diabetes can promote tissue Ang II/AT1R actions. Firstly, hyperglycemia directly stimulates local Ang II production in cardiomyocytes [163], cardiac fibroblasts [164], and endothelial cells [165] as well as in murine and human diabetic heart tissues [164, 166]. For example, Fiordaliso and colleagues demonstrated that glucose increased Ang II in cardiac myocytes, which made them more susceptible to undergo apoptosis [167], consistent with the view that the RAAS promotes the development of diabetic cardiomyopathy [162, 166]. Studies on cardiac myocytes suggest that the mechanism whereby hyperglycemia increases local Ang II in the heart is the generation of intracellular Ang II by intracellular chymase and/or internalized prorenin. Then, intracellular Ang II could directly produce oxidative stress and cellular apoptosis and/or enhance RAAS components expression through a positive feedback mechanism [168].

The second mechanism underlying AngII/AT1R activation in diabetes is that high glucose concentrations can enhance the tissue response to Ang II and vice versa. While Ang II can increase the activation of glucose-induced transcription factors in vascular smooth muscle cells [169], hyperglycemia can increase the contractile aortic response to Ang II [170]. This additive effect could be ascribed to the fact that diabetes induces AT1R expression in the heart [171, 172] and the vasculature [172]. Interestingly, several works have demonstrated that hyperglycemia can also stimulate aldosterone secretion by increasing local Ang II. Xue and Siragy observed an upregulation of renal aldosterone synthase in diabetic rats, which was significantly reduced by AT1R blockade [173]. Similar findings were reported in podocytes [174] and in cardiomyocytes [175]. Locally, aldosterone triggers a vicious cycle that promotes AngII/AT1R effects as it mediates part of Ang II effects [176], increases tissue ACE and AT1R [177], and reduces tissue ACE2 [178, 179]. Moreover, together with Ang II, aldosterone promotes oxidative stress, fibrosis, and apoptosis. Therefore, many studies are now supporting at least a synergistic role for aldosterone in the pathogenesis of diabetic cardiovascular complications [180].

The third way whereby diabetes promotes Ang II tissue actions is through the several metabolic abnormalities associated with hyperglycemia. These include advanced glycation end products, which form after prolonged hyperglycemia and oxidative stress, dyslipidemia, and low-grade inflammation. All of them can in fact stimulate the AngII/AT1R pathway by upregulating AT1R expression [181–184]. An interesting aspect that should be considered is that, in the heart, the conversion of Ang I to Ang II relies not only on ACE activity but also on other enzymes such as tissue chymase [185, 186], which are found primarily in mast cells. Given this localization, chymase might link inflammation to RAAS activation and CVD. Unfortunately, inhibition of this chymase is yet to be realized [187].

Another intriguing mechanism whereby diabetes enhances AngII/AT1R actions is ACE2 downregulation, which does not only promote Ang II actions but also reduce local Ang 1–7 leading to an imbalance of the RAAS. This concept is supported by the works of Tikellis and colleagues, who showed that the induction of diabetes was associated with a significant reduction of ACE2 expression and ACE2 activity in the heart and the vasculature [172] together with a significant increase in circulating Ang II and a significant reduction of Ang 1–7 levels [172]. Similar changes were reported in the kidney of diabetic mice [132, 188]. It has been argued that ACE2 may be more important than ACE in regulating cardiac levels of Ang II and Ang 1–7 and therefore more important for balancing RAAS activation [189]. This concept relies on the experimental evidence that while ACE2 deficiency results in increased cardiac levels of Ang II [51] and reduced levels of Ang 1–7 [132], ACE deficiency does not modify cardiac Ang II, which could be generated by non-ACE pathways [190], including chymases.

Therefore, ACE2 replenishing strategies could be an important therapeutic tool against diabetic CVD. Another promising therapeutic strategy against diabetic CVD seems to be AT2R stimulation. The rationale behind it is that diabetes seems to upregulate AT2R expression [24], AT2R expression was found increased in the hearts of diabetic rats [171, 191], as well as in their kidneys [192]. Similar results were obtained in kidney biopsies of patients with type 2 diabetes [193]; this is in line with the report that glucose induced AT2R renal expression both in vivo [194, 195] and in vitro through the expression of the transcriptor factor IRF-1 [195].

Lastly, diabetic patients have higher prorenin levels than normal subjects, and their prorenin levels are 10-fold
higher than their renin levels [196]. Given that the binding of prorenin to (P)RR can generate Ang I and also stimulate (P)RR intracellular signaling, it has been argued that plasma prorenin could contribute more substantially than renin to the pathogenesis of end-organ damage [196]. This is supported by the finding that diabetic patients with high prorenin levels had a higher risk of microvascular complications [197], whose development could be predicted by the same prorenin levels [198]. Consistent with these circulating changes, Connelly and colleagues demonstrated that, also at a tissue level, diabetic cardiomyopathy was associated with a 3-fold increase in both (P)RR gene and protein expression, which were reduced by the renin inhibitor Aliskiren. Although the increased expression of (P)RR in the diabetic heart was interpreted as a beneficial response to limit intracellular acidosis, due to its sequence identity with the V-ATPase, it was argued that (P)RR abundance could still modulate the activity of the local RAAS and expose cardiac myocytes to Ang II increased concentrations [199].

4. New Potential Therapies against Diabetic Cardiovascular Complications

4.1. New Agents for Classic Targets

4.1.1. Angiotensin Receptor-Neprilysin Inhibition. ACE inhibition, AT1R antagonism, and MR blockade are some of the most classic therapeutic strategies against CVD. Another classic therapeutic strategy against heart failure is to increase natriuretic peptide levels, as they are natriuretic, diuretic, vasodilating, and able to inhibit pathologic growth in heart failure [200]. However, although the idea to boost their activity in patients with heart failure seemed theoretically infallible, the approaches used for years have produced only partial if not disappointing effects. These approaches included short-term intravenous infusions of natriuretic peptides or inhibition of neprilysin, which is the enzyme that degrades natriuretic peptides, together with bradykinin, adrenomedullin, and possibly other substrates such as Ang II.

So, the most likely explanation underlying the disappointing effect of the oral inhibitor of neprilysin was that neprilysin degraded also Ang II, and therefore inhibiting neprilysin would increase not only natriuretic peptides but also Ang II actions. Eventually, the combination of an ACE and a neprilysin inhibitor solved this problem and turned out to be superior to either approach in terms of cardiovascular benefits [201, 202]. Unfortunately, in clinical trials this combination was associated with angioedema [203, 204]. In order to overcome this issue, angiotensin receptor-neprilysin inhibitors (ARNis) have been developed, such as LCZ696, which is an association of the ARB Valsartan with the neprilysin inhibitor Sacubitril (Figure 4). In a recent trial this association was found superior to Enalapril in reducing the risk of death and hospitalization in patients with heart failure [205], including those with diabetes [206]. This is consistent with the observation that the administration of LCZ696 in an experimental model of diabetes significantly reduced cardiac hypertrophy and fibrosis and improved ejection fraction after myocardial reperfusion injury [207].

4.1.2. Aldosterone Synthase Inhibition. Another classic target against CVD is aldosterone, due to the growing appreciation of its contribution to CVD development and progression. The drugs that are available for blocking aldosterone actions are Spironolactone and Eplerenone, which are both mineralocorticoid-receptor antagonists. Nevertheless, Spironolactone and Eplerenone have still limited clinical use, due to the poor selectivity of Spironolactone, the low potency of Eplerenone, and the fact that only MR-dependent effects of aldosterone can be inhibited. An alternative approach to antagonize aldosterone is inhibiting its formation. For this reason, several aldosterone synthase (CYP11B2) inhibitors have been developed. The compounds targeting CYP11B2 that have been studied so far are FAD286, which has been used mostly in experimental settings, and LC1699, which has been developed for human use in 2010 [208]. A few works have demonstrated that FAD286 reduced mortality, improved cardiac remodeling [209, 210], and reduced atherosclerosis [211]. Nevertheless, both drugs have exhibited clear side effects due to the inhibition of CYP11B1, which synthesizes glucocorticoids. Moreover, another concern prior to their use is that aldosterone synthase inhibitors could allow the activation of unprotected MR by glucocorticoids, and this needs further research.

4.1.3. Renin Inhibition. Renin inhibition represents another very attractive therapeutic strategy to reduce Ang II levels. This can be achieved with the renin inhibitor Aliskiren, which attenuates Ang II generation and reduces blood pressure comparably to ACE inhibitors and ARB [212]. However, similar to what was reported in the ONTARGET trial, the ALTITUDE study has demonstrated that the simultaneous administration of Aliskiren with an ACE inhibitor or an ARB should be avoided [213]. A critical concern on the use of Aliskiren is that it is associated with high renin levels that might bind to (P)RR [214, 215]. Therefore, specific blockade of (P)RR could provide more benefits as compared to renin inhibition alone, as it could not only reduce (pro)renin enzymatic activity but also prevent some AngII-independent effects of (pro)renin. So far, a (P)RR antagonist, called HRP (“handle region” peptide), has been designed. In particular, this molecule mimics the “handle region” of prorenin, which allows its binding to (P)RR and its subsequent nonproteolytic activation. Studies in animal models of diabetes show promising results as reviewed in [148], possibly because HRP exerts its effects only in diseases associated with high prorenin and low renin levels, such as diabetes.

4.2. New Agents for New Targets

4.2.1. ACE2 Replenishing Strategies. A promising therapeutic strategy in cardiovascular medicine is represented by RAAS modulation. As compared to RAAS antagonism, RAAS modulation combines ACE/AngII/AT1R blockade with the stimulation of ACE2/Ang 1-7/Mas1R and AT2R. The latter can be achieved by a series of new therapies that includes ACE2 replenishing strategies, Ang 1–7 administration, and AT2R agonists. However, also ACE inhibitors and AT1R blockers should be considered RAAS modulating agents,
as they can increase ACE2 expression and Ang 1–7 levels severalfold, suggesting that part of their beneficial effects is due to ACE2/Ang 1–7 effects [216, 217].

Having said that, Crakower and colleagues showed that ACE2 is an essential regulator of cardiac function, by demonstrating that ACE2 deficiency was associated with reduced systolic function [218]. Similar findings have been reported in animal models of type 1 and type 2 diabetes [219, 220]. In addition to that, we have demonstrated that ACE2 deficiency was associated with accelerated atherosclerosis [51]. It has also to be noted that ACE2 deficiency might in fact influence not only the risk of developing CVD but also the response to the treatment, particularly in the setting of diabetes where ACE2 expression is constitutively low [188]. These findings emphasize the potential utility of ACE2 repletion as a strategy to reduce CVD. Current therapeutic tools that modulate ACE2 levels or activity, which are still under investigation, include adenoviral ACE2 gene transfer, recombinant human ACE2 (rhACE2), ACE2 activators, oral ACE2, and Ang 1–7 bioencapsulated in plant cells. So far, both ACE2 gene transfer and the administration of an ACE2 activator have ameliorated diabetic cardiomyopathy [221, 222]. Moreover, the reports that rhACE2 administered intravenously to healthy human subjects was well tolerated [223] and that it resulted in sustained reduction in plasma Ang II levels and elevation in Ang 1–7 levels [131] are encouraging, as they are shortening the gap between bench and bedside with respect to the possibility of using rhACE2 in clinical practice.

4.2.2. Ang 1–7 Administration. Based on the protective cardiovascular effects of Ang 1–7, several experimental studies have tested the hypothesis that Ang 1–7 infusion could ameliorate diabetic cardiomyopathy. In all these works, Ang 1–7 improved all the structural hallmarks of diabetic cardiomyopathy, which is characterized by left ventricle hypertrophy and left and right ventricle fibrosis and dysfunction [224–226]. In db/db mice with early [227] and advanced disease [228], Ang 1–7 reversed diabetes-induced changes. In particular, in the early disease model [227], Ang 1–7 increased cardiac output and cardiac index and decreased cardiomyocyte hypertrophy, heart fibrosis, and inflammation. Vascularization was also improved, which correlated with increased numbers of bone marrow residing in and circulating endothelial and mesenchymal stem cells [227]. Here, Mas1R seemed to be the main receptor mediating Ang 1–7 effects, which explains why AVE0991, which is a Mas1R agonist, had cardioprotective effects in diabetic rats [229]. In addition, in the advanced disease model, Ang 1–7 ameliorated diastolic function and decreased not only cardiac hypertrophy and fibrosis but also triacylglycerol accumulation [228]. Moreover, consistent with the report that Ang 1–7 preserved cardiac function, coronary perfusion, and aortic

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**Figure 4: Mechanisms of action of angiotensin receptor/neprilysin inhibitors.** ARB antagonize Ang II binding to AT1R, which causes hypertension and multiorgan injury. NEPi inhibit the activity of NEP, which is an enzyme degrading natriuretic peptide, bradykinin, and other peptides. The result is a reduction of Ang II harmful effects and an increase of the benefits of natriuretic peptides and bradykinin. ACE is for angiotensin-converting enzyme; ADM is for adrenomedullin; Ang is for angiotensin; ANP is for atrial natriuretic peptide; ARB is for angiotensin II receptor blockers; AT1R is for Ang II type 1 receptor; BNP is for brain natriuretic peptide; NEPi is for neprilysin inhibitors.
endothelial function in a rat model of heart failure [230]. Ang 1–7 improved cardiac recovery from ischemia/reperfusion and restored the normal reactivity to constrictor and dilator stimuli in the vasculature [231]. Recently, we have shown that Ang 1–7 administration significantly reduced ex vivo leukocyte recruitment in an animal model of diabetes. This was associated with a reduction of glucose-induced molecule adhesion expression and leukocyte adhesion to endothelial cells in vitro [232]. These effects were completely blocked by the MasR antagonist, suggesting that MasR was the main receptor mediating Ang 1–7 effects on endothelial cells.

4.2.3. AT2R Agonists. AT2R activation has been currently achieved by compound 21 (C21), which is a nonapeptide that acts as a highly selective AT2R agonist and stimulates AT2 receptors. Several studies have shown its efficacy in reducing tissue fibrosis in the cardiovascular system. However, those that have been carried out in diabetic conditions have explored its actions mostly against renal fibrosis. In this setting, C21 was able to significantly reduce renal fibrosis in experimental models of both type 1 [233] and type 2 diabetes [234]. In one of these studies, C21 was able to significantly reduce also the expression of several inflammatory mediators [233]. This finding is consistent with the vascular effects of this drug, which prevented endothelial inflammation and reduced leukocyte adhesion both in vivo and in vitro [235]. In addition, recently, C21 was found to be able to reduce diabetes-associated atherosclerosis [236].

5. Conclusions
The RAAS has been studied for more than a century. Nevertheless, the current picture of the RAAS is that of an extremely complex pathway, which has not been fully characterized yet and might hold in store new aspects that have still to be discovered. Certainly, what we do know is that blocking Ang II reduces cardiovascular complications. This is particularly true in diabetic conditions, where Ang II/AT1R pathway is activated, while the Ang 1–7/Mas1R is not. Therefore, the aim of new therapies is not only to block Ang II harmful effects but also to augment the activity and actions of potentially beneficial pathways, by ACE2 replenishing strategies, Ang1–7 administration, and AT2R agonists. Here is another paradigm shift: treatment of diabetic cardiovascular disease should move from RAAS inhibition to RAAS modulation.

Competing Interests
The authors declare that there are no competing interests regarding the publication of this article.

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