Angiotensin receptor blockers for the reduction of proteinuria in diabetic patients with overt nephropathy: results from the AMADEO study

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Abstract: Diabetic kidney disease is characterized by persistent albuminuria (>300 mg/dl or >200 μg/min) that is confirmed on at least 2 occasions 3 to 6 months apart, with a progressive decline in the glomerular filtration rate (GFR), elevated arterial blood pressure, and an increased risk for cardiovascular morbidity and mortality. Diabetic kidney disease is the leading cause of end stage renal disease (ESRD) prompting investigators to evaluate mechanisms by which to slow disease progression. One such mechanism is to block the activity of angiotensin II at the receptor site and agents that follow this mechanism are referred to as angiotensin receptor blockers (ARB). There is sufficient clinical evidence to support that ARB have protective effects on kidney function in patients with diabetes and hypertension. However, in the past decade there have been few investigations comparing individual ARBs on renal outcomes. Telmisartan, a lipophilic ARB with a long half-life, has been hypothesized to have a greater anti-proteinuric effect when compared to the shorter acting losartan. Therefore, the A comparison of telMisartan versus losArtan in hypertensive type 2 DiabEtic patients with Overt nephropathy (AMADEO) trial sought to investigate renal and cardiovascular endpoints. In this review, we discuss the pathophysiology of diabetic kidney disease and implications of the AMADEO trial in the context of current understanding from recent outcome trials.

Keywords: diabetic kidney disease, hypertension, telmisartan, AMADEO

Introduction

Diabetic kidney disease (DKD) is the leading cause of end stage renal disease (ESRD) and is a multifactorial combination of hemodynamic and metabolic abnormalities that collectively contribute to kidney damage resulting in proteinuria and reductions in glomerular filtration rate (GFR). Recent data support that proteinuria is a surrogate maker for cardiovascular risk and reductions in proteinuria correlate with declines in cardiovascular morbidity and mortality. Thereby, interventions that target blood pressure control and proteinuria, specifically interruption of the renin-angiotensin system (RAS) with either angiotensin converting enzyme (ACE) inhibitors or angiotensin II receptor blockers (ARB), have been utilized in attenuating the progression of DKD. Among available ARBs, telmisartan has been reported to have a greater lipophilicity, longer half-life, and debatably the most consistent reductions in blood pressure. Therefore, investigators recently sought to compare telmisartan with losartan, which has less lipophilicity and a shorter duration of action, in patients who had overt DKD (urinary protein to creatinine ratio ≥ 700) in the A comparison of telMisartan versus losArtan in hypertensive type 2 DiabEtic patients with Overt nephropathy (AMADEO) trial. Investigators reported that telmisartan was superior to losartan in reducing proteinuria in hypertensive patients with DKD with relatively similar reductions in blood pressures. Further, the authors proposed that the superiority of telmisartan could
be due to its intrinsic peroxisome proliferator-activated receptor–gamma (PPAR-γ) agonist properties.

The incidence of DKD continues to increase in the US and globally. Understanding the mechanisms underlying the development of DKD is essential for establishing novel therapeutic strategies for the prevention or arrest of progressive disease. Herein, we will review some of these mechanisms as they relate to the AMADEO trial findings.

**Pathophysiology and markers of diabetic kidney disease**

Evidence suggests that up to 44% of patients with diabetes mellitus develop DKD. Development of DKD is associated with progressive functional and structural changes in the basic kidney unit, ie, the nephron and glomerulus, affected via hemodynamic and metabolic pathways. Hemodynamic and metabolic factors contribute equally towards the development of DKD, it is now clear that these processes are interlinked. Earlier stages of DKD include a hyperfiltration mechanism that occurs due to decreased resistance of both afferent and efferent arteriole. Afferent arteriole has greater decrease in resistance than its efferent counterpart. There is defective autoregulation of tone due to complex interaction of mediators including prostanoids, nitric oxide, reactive oxygen species (ROS), lipids, vascular endothelial growth factor (VEGF), transforming growth factor-beta 1 (TGF-β1), high glucose and the RAAS, specifically angiotensin II (Ang II). These hemodynamic changes and the defect in autoregulation allow an increased filtration of albumin at the level of the glomerulus. It has been shown that proteinuria can occur as a result of molecular and structural abnormalities in the podocyte slit diaphragm within the glomerular epithelial cell. Ang II has been reported to be a primary mediator of loss of the slit-pore diaphragm. In addition to promoting glomerular nephrin depletion, Ang II also appears to have other actions that promote the development of proteinuria, including trophic effects on the kidney and increasing glomerular membrane pore size. This in turn promote structural changes like mesangial cell proliferation, thickening of basement membrane that further potentiate injury to podocytes. At the molecular level, hyperglycemia and proteins altered by high blood glucose such as Amadori products and advanced glycation end products (AGEs) are key players in the development of DKD. Evidence suggests that an increase in reactive oxygen species (ROS) formation induced by high glucose-mediated activation of the mitochondrial electron-transport chain is an early event in the development of diabetic complications. A variety of tissue growth factors and cytokines are then induced through complex signal transduction pathways involving protein kinase C (PKC), mitogen-activated protein kinases (MAPK), and the transcription factor NF-κB. High glucose, AGEs, and ROS act in concert to induce growth factors and cytokines. Particularly, tissue growth factors are important in the development of renal hypertrophy and accumulation of extracellular matrix components.

Activation of the renin-angiotensin-aldosterone system (RAAS) by hemodynamic changes, high glucose, mechanical stress, and proteinuria with an increase in local formation of Ang II contribute many of the pathophysiological changes associated with DKD (Figure 1). In fact, it has been shown that Ang II is involved in almost every pathophysiological process implicated in the development of DKD (hemodynamic changes, hypertrophy, extracellular matrix accumulation, growth factor/cytokine induction, ROS formation, podocyte damage, proteinuria, interstitial inflammation, adverse effects on the balance between the fibrinolytic and thrombotic mechanisms). Therefore, interruption of the RAAS via inhibition of Ang II is an essential part of every therapeutic regimen in managing the patient with DKD.

**Role of drugs blocking RAAS**

RAAS inhibition is the mainstay of therapy in preventing the progression of DKD. In addition to normalization of systemic and glomerular hypertension, it is now clear that inhibition of the RAAS at various levels by the Ang II antagonists (ACE inhibitors and ARBs) have several renoprotective effects, including anti-inflammatory and anti-fibrotic mechanisms. Addition of diuretics has been strongly recommended by KDOQI guidelines to achieve target blood pressures of ≤130 mmHg in cases of recalcitrant hypertension in DKD patients. The role for aldosterone antagonist (spironolactone), renin antagonist (aliskiren) and statins in prevention of metabolic derangements consequent to RAAS overactivity in DKD is emerging. In whole animal models, experiments to tease out the non-hemodynamic renoprotective effects of the RAAS antagonists included transgenic rats with a human AT₁ receptor (AT₁ R) gene which overexpressed AT₁ R. These rats developed significant albuminuria, podocyte effacement, progressive fibrosis, leading to focal segmental glomerulosclerosis (FSGS), which were independent of blood pressure and reversed by the blockade of Ang II. Aldosterone has been shown to cause kidney oxidative stress, inflammation, fibrosis, and sclerosis. Shibata et al demonstrated that aldosterone infusions and high-salt diet, when administered to uninephrectomized normotensive
Sprague Dawley (SD) rats, led to glomerulosclerosis, hypertension, and kidney damage. Administration of the mineralocorticoid receptor (MR) blocker (spironolactone) ameliorated and reversed the harmful effects of aldosterone. A recent study, Aliskiren in the EValuation of PrOteinuria In Diabetes (AVOID) Trial, showed that the addition of a renin inhibitor to an ARB in diabetic and hypertensive patients decreases proteinuria further when compared with an ARB alone. Statins have been proposed to have pleiotropic effects, actions independent of their cholesterol-lowering mechanism. These include decreased ROS generation via inhibition of NADPH oxidase and therefore have a role as renoprotective agent. Large randomized controlled trials designed to target DKD populations may be able to provide comprehensive answers as to the usefulness of these drugs.

Strong evidence points to the expression of all RAAS components in the kidneys, including the podocytes and tubular cells. High glucose stimulates expression of renin and angiotensinogen in mesangial and tubular cells. This stimulation results in an increase in local Ang II concentrations which may, in turn, through autocrine and paracrine pathways induce several cytokines and growth factors. Therefore blockade of the overexpressed RAAS will lead to renoprotection. Taking into account the many effects of the RAAS in the pathophysiology of DKD, early treatment with drugs interfering with the tissue effects of RAAS are a necessary prerequisite to prevent development and progression of DKD.

**Lessons from clinical trials on RAAS-blockade**

Blockade of the RAAS is pivotal in the management of DKD as described above. Strong evidence favors the use of ACE inhibitors and ARBs in patients with type 2 diabetes.
**Table 1** Recent clinical trials leading up to the AMADEO study

<table>
<thead>
<tr>
<th>Trial/year</th>
<th>Patient population</th>
<th>N</th>
<th>Treatment arms</th>
<th>Primary endpoint</th>
<th>Final outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOPE 1999</td>
<td>High-risk patients without known diabetes</td>
<td>5720</td>
<td>Ramipril/placebo</td>
<td>New onset overt nephropathy</td>
<td>22% Reduction in new onset overt nephropathy (RR 0.66; 95% CI 0.51–0.85; p &lt; 0.001).</td>
</tr>
<tr>
<td>IDNT 2001</td>
<td>Hypertension with diabetic nephropathy</td>
<td>1715</td>
<td>Irbesartan/amlodipine/placebo</td>
<td>Composite of development of ESRD, doubling Cr.</td>
<td>Irbesartan significantly reduced primary composite end point by 20% compared to placebo (p &lt; 0.02) and by 23% compared to amlodipine (p &lt; 0.006).</td>
</tr>
<tr>
<td>IRMA 2001</td>
<td>Type 2 diabetes with MA</td>
<td>590</td>
<td>Irbesartan/placebo</td>
<td>Time to onset of diabetic nephropathy</td>
<td>Irbesartan reduced risk of primary end point (HR 0.30; 95% CI 0.14–0.61; p &lt; 0.001 for 300 mg irbesartan; HR 0.61; 95% CI 0.34–0.99; p &lt; 0.08 for 150 mg irbesartan).</td>
</tr>
<tr>
<td>RENAAL 2001</td>
<td>Type 2 diabetes with nephropathy</td>
<td>1513</td>
<td>Losartan/placebo</td>
<td>Composite of doubling of Cr, development of ESRD.</td>
<td>Losartan significantly reduced the composite end point by 16% (p &lt; 0.02), but not death.</td>
</tr>
<tr>
<td>MARVAL 2002</td>
<td>Type 2 diabetes with MA</td>
<td>332</td>
<td>Valsartan/amlodipine</td>
<td>% change in UAER from BL to 24 Wks.</td>
<td>UAER at 24 wk was reduced by 44% with valsartan and 8% with amlodipine (p &lt; 0.001); valsartan significantly reversed MA to normal albuminuria.</td>
</tr>
<tr>
<td>BENEDICT 2004</td>
<td>Type 2 diabetes and HTN but with normo-albuminuria</td>
<td>1204</td>
<td>Trandlopril/verapamil/placebo</td>
<td>Development of persistent MA</td>
<td>For developing end point AF = 0.39 for trandlopril + verapamil vs placebo. AF = 0.44 for trandlopril vs placebo and AF = 0.83 with verapamil vs placebo.</td>
</tr>
<tr>
<td>DETAIL 2004</td>
<td>Type 2 diabetes with early nephropathy</td>
<td>250</td>
<td>Telmisartan/enalapril</td>
<td>Change in the GFR (determined by measuring the plasma clearance of iohexol)</td>
<td>Change in the GFR was −17.9 mL/min/1.73 m² of body-surface area with telmisartan, as compared with −14.9 mL/min/1.73 m² with enalapril, for a treatment difference of −3.0 mL/min/1.73 m² (95% CI −7.6–1.6 mL/min/1.73 m²).</td>
</tr>
<tr>
<td>TRENDSY 2007</td>
<td>Type 2 diabetes, HTN, GFR &gt; 80 mL/min, and normo- or micro-albuminuria</td>
<td>96</td>
<td>Telmisartan/ramipril</td>
<td>Increase in renal plasma flow (RPF)</td>
<td>Telmisartan increased RPF from 652.0 ± 27.0 to 696.1 ± 31.0 mL/min (p = 0.047), whereas ramipril produced no significant changes in RPF (r = 0.47, p &lt; 0.001).</td>
</tr>
<tr>
<td>INNOVATION 2007</td>
<td>Type 2 diabetes and UACR 100-300mg/g, Cr &lt; 1.5 mg/dl (men) and &lt; 1.3 mg/dl (women).</td>
<td>527</td>
<td>Telmisartan (80/40 mg)/placebo</td>
<td>Transition rate incipient to overt Nephropathy, UACR &gt; 300 mg/g, and increase ≥ 30% from baseline at 2 consecutive 4-week visits.</td>
<td>Transition rates to overt nephropathy were 16.7% with 80 mg telmisartan (n = 168), 22.6% with 40 mg telmisartan (n = 172), and 49.9% with placebo (n = 174) (both telmisartan doses/ placebo, p &lt; 0.0001).</td>
</tr>
<tr>
<td>VIVALDI 2008</td>
<td>HTN SBP/DBP &gt; 130/80 mmHg patients with type 2 diabetes, proteinuria (≥900 mg/24 h) and Cr (≥3.0 mg/dL)</td>
<td>885</td>
<td>Telmisartan/valsartan</td>
<td>Change from BL of 24 h proteinuria.</td>
<td>Comparable reduction in 24 h urinary protein excretion rates. Geometric mean reduction (95% confidence interval) telmisartan, 33% (27%–39%); valsartan, 33% (27%–38%).</td>
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(Continued)
and overt nephropathy. Current management consensus statements from the National Kidney Foundation (NKF)/Kidney Disease Outcomes Quality Initiative (KDOQI), American Diabetes Association (ADA), and Joint National Committee (JNC) VII advocate the use of ARBs for management of patients with hypertension and DKD, even in those with advanced stages of nephropathy as well as in those with microalbuminuria. ARBs have been shown to exert their beneficial effect by blood pressure reduction, improving endothelial dysfunction and reducing oxidative stress.

Data from the Heart Outcomes Prevention Evaluation (HOPE) Study evaluated microalbuminuria as a progressive manifestation of overt type nephropathy in patients with type 2 diabetes as a secondary endpoint. Ramipril was shown to reduce the onset of overt nephropathy by 22%. Similar observations were reported in the Irbesartan type II Diabetic Nephropathy Trial (IDNT), irbesartan was superior to both placebo and amlodipine in improving time to the primary endpoint, which was the composite of doubling of baseline serum creatinine and development of ESRD. Interestingly, the benefits of irbesartan were seen with similar reductions in blood pressure when compared to amlodipine, revealing non-hemodynamic effects of ARBs. However, this trial did not include ACE inhibition in its treatment arms. Another caveat that was borne out of post-hoc analyses was that initial changes in proteinuria were not independent predictors of long-term renal outcomes, but were dependent on specific treatment arms (in this case irbesartan). This effect was later tested in the AMADEO trial. Analysis of IDNT showed that over a follow up period of 2.6 years there was a 2.2 times increase in risk of doubling serum creatinine or ESRD in patients with systolic blood pressure (SBP) > 149 mmHg compared with those with SBP < 143 mmHg regardless of the intervention used for treatment of blood pressure. Among the patients achieving comparable reduction in SBP, the incidence of renal endpoints was lower in patients treated with irbesartan compared to amlodipine.

The Irbesartan in Patients with Type 2 Diabetes and Microalbuminuria Study (IRMA) further strengthened the results of IDNT by showing that irbesartan has a dose-dependent improvement in time to onset of DKD. After a follow up of 2 years, it was found that only 5.2% patients treated with irbesartan 300 mg progressed to overt nephropathy as opposed to 9.7% of those receiving irbesartan 150 mg and 14.9% of those receiving placebo. In fact, the IRMA study group hypothesized that the renoprotective effect of irbesartan was not fully evident in their original study due to dose limitations and they demonstrated even greater renal protection with 900 mg of irbesartan and minimal increase in side effects (IRMA-2). The Reduction of Endpoints in NIDDM with the Angiotensin II Antagonist Losartan Study (RENAAL) study was similar to the IDNT and IRMA studies and demonstrated delayed time to onset of renal dysfunction in the losartan treated group. Post hoc analyses of proteinuria as secondary endpoint showed that early response to losartan therapy is an important predictor of long term renal protection. Both baseline and residual (6 month) albuminuria were linearly correlated with renal outcomes. The authors recommended reduction in albuminuria to the lowest achievable level as the optimum goal for therapy.

MicroAlbuminuria Reduction With VALsartan (MARVAL) Study targeted urine microalbumin excretion (UAER) as the primary endpoint and demonstrated reduction in albuminuria to 56% of its baseline in both normotensive and hypertensive
type 2 diabetics after 6 months of intention to treat therapy.\textsuperscript{25} Importantly, the improvement in albuminuria was independent of blood pressure, as the amlodipine treatment arm achieved similar reductions in blood pressure, once again demonstrating non-hemodynamic effects of ARBs in modulating pathophysiology of DKD.

The Bergamo Nephrologic Diabetes Complications Trial (BENEDICT) compared the efficacy of ACE inhibitor trandolapril versus non-dihydropyridine calcium channel blocker verapamil versus trandolapril alone versus verapamil alone versus placebo in delaying the incidence of microalbuminuria in type 2 diabetics.\textsuperscript{26,27} The trandolapril containing arms demonstrated significant delay in the development of microalbuminuria. Interestingly, ONTARGET evaluated the use of ACE inhibitor in prevention of early chronic kidney disease.\textsuperscript{28} RAAS control with ACE inhibitor has been shown to prevent the emergence of persistent microalbuminuria in patients with type 2 diabetes and apparently normal urinary albumin levels (<20 μg/min).

The Diabetics Exposed to Telmisartan and Enalapril (DETAIL) study was among the first studies to compare the efficacy of long-acting telmisartan with ACE inhibitor enalapril.\textsuperscript{29} Although there was an initial advantage in slowing down the decline in GFR with telmisartan, end of trial analyses could only demonstrate non-inferiority over enalapril. Based on the findings of DETAIL, it may be argued that a more aggressive approach to renoprotection may be required in the long term.\textsuperscript{18} The recently published Telmisartan versus Ramipril in renal Endothelial DYsfuntion (TRENDY) study provided the first head-to-head comparison of an ARB and an ACE inhibitor on renal endothelial function.\textsuperscript{30} TRENDY was performed in 96 patients with type 2 diabetes who had mild-to-moderate hypertension (seated SBP/diastolic blood pressure [DBP] 140 to 180/90 to 110 mm Hg), normo- or microalbuminuria, and GFR > 80 mL/minute/1.73 m². Patients were randomized to either telmisartan 40 mg or ramipril 5 mg for 3 weeks, followed by telmisartan 80 mg or ramipril 10 mg for 6 weeks. Both telmisartan and ramipril improved nitric oxide activity (a measure of endothelial function), with a trend toward superiority of telmisartan. There was also a significant decrease in urinary albumin excretion from 9.0 to 7.3 mg/24 hours with telmisartan (p = 0.022), whereas the change from baseline with ramipril was not significant.

In the study to inVestigate the efficacy of telmiSartan versus VALsartan in hypertensive type 2 Diabetic patients with overt nephropathy (VIVALDI), both telmisartan and valsartan improved proteinuria by 33% at the end of 1 year.\textsuperscript{31} However, the benefits from telmisartan were not significantly greater than valsartan in this intention to treat analysis. The authors did note that patients in the valsartan treatment arm tended to receive additional antihypertensives in a greater proportion than the telmisartan group most probably due to the shorter half-life of the former drug.\textsuperscript{31} Moreover, proteinuria measurements were based on 24-hour urine protein collections. The INcipient to OVert: Angiotensin II receptor blocker, Telmisartan, Investigation On type 2 diabetic Nephropathy (INNOVATION) trial showed that telmisartan is able to prevent the transition to overt renal disease in Japanese patients after a mean follow-up of 1.3 years.\textsuperscript{32} The relative risk of transition to overt nephropathy was reduced by 66% and 55%, respectively, with telmisartan 80 mg and telmisartan 40 mg compared with placebo in the overall patient population. Thus, for every 3 patients treated, telmisartan 80 mg prevented 1 transition to overt nephropathy over the course of approximately 1 year. The renoprotective benefit of telmisartan was also apparent in patients who were normotensive at baseline, which suggests that there is a blood pressure-independent effect. In addition, INNOVATION showed treatment of patients with telmisartan could bring about regression of incipient disease, with 21.2% and 12.8% of patients in the telmisartan 80 mg and 40 mg arms respectively, becoming normoalbuminuric compared with 1.2% of patients receiving placebo (both doses p < 0.001).\textsuperscript{32} Interestingly, both the DROP trial and the IRMA-2 substudy demonstrated a greater anti-proteinuric effect with ultra-high doses of ARBs while not significantly achieving any further lowering in blood pressure.\textsuperscript{33,34,35}
itself cannot be taken as a definitive marker of improved renal function, and that the benefits of any treatment, including combination blockade of the RAS on major renal outcomes, remain to be elucidated.

**Rationale for AMADEO trial**

Substantial evidence has accumulated over the last decade in favor of using RAAS inhibitors for slowing down the progression and onset of DKD. Use of ARB and ACE inhibition is clearly superior to the use of non-RAAS blood pressure control medications in ameliorating renal damage at similar levels of blood pressure control. This may be due to the non-hemodynamic and metabolic improvements in the cardiovascular system including better endothelial function, reduced oxidative stress, decrease in inflammation and fibrosis. Evidence has emerged demonstrating that a >30% reduction in proteinuria at 6 months to 1 year following initiation of treatment was associated with slower progression of DKD and a reduction in the incidence of cardiovascular events, independent of reductions in blood pressures. The ARBs have been shown to be non-inferior to ACE inhibitors in achieving blood pressure control and reductions in proteinuria, often leading to interchangeable regimens based on these drugs depending on the tolerability by patients.

Telmisartan, an ARB with high lipophilicity, a high level of angiotensin II type-1 (AT_1_) receptor (AT_1_ R) binding and a long half-life is currently FDA approved for the treatment of hypertension but is not indicated yet for the treatment of proteinuria in patients with diabetes. Instead, losartan is currently approved as a first-line treatment for DKD. Telmisartan has been reported to achieve smoother blood pressure control with fewer side effects when compared to other ARBs. In addition, animal experiments have demonstrated other potential mechanisms for its beneficial effects including PPAR-γ agonist properties. All of these properties together may make telmisartan an ideal drug to achieve the target of >30% reduction in proteinuria ultimately resulting in a slower progression to kidney failure.

**Pharmacology of telmisartan**

Telmisartan is a potent, long-lasting, non-peptide antagonist of the AT_1_ receptor. It selectively and insurmountably inhibits stimulation of the AT_1_ receptor by angiotensin II without affecting other receptor systems involved in cardiovascular regulation. Very high lipophilicity, a unique feature of telmisartan, coupled with a high level of distribution; indicate that the compound offers a clinically important advantage of good tissue penetration (Table 2). Due to its physicochemical properties, telmisartan shows excellent oral absorption and tissue penetration.

**Affinity for AT_1_ receptor**

Studies have demonstrated that telmisartan has the strongest binding affinity to AT_1_ receptor among various ARBs (Table 2). The rank order of affinity is in the following order; telmisartan > olmesartan > candesartan > EXP3174 > valsartan > losartan. The findings of AMADEO suggest that telmisartan may have long-lasting blood pressure-lowering effects and superior cardioprotective properties in patients with hypertension due to its strong AT_1_ receptor antagonistic ability.

**Pharmacokinetics**

Telmisartan is not a pro-drug and has a longer terminal elimination half-life than other commercially available sartans (~24 hours), making it suitable for once-daily dosing. The compound is not metabolized by cytochrome P450 isoenzymes and has a low risk for P450-based drug interactions. A relevant fact to be noted is that unlike other ARBs, where up to 50% of the dose is excreted by the kidney, only 1% of a telmisartan dose in excreted by the kidney. Therefore renal impairment is unlikely to affect the pharmacokinetics of telmisartan. Telmisartan has a pharmacokinetic profile that allows differentiating it from other angiotensin receptor blockers; long duration of action (mean terminal half-life of 24 hours, the longest in its class) and high tissue penetration (illustrated by the

**Table 2 Pharmacokinetics of ARBs**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Bioavailability (%)</th>
<th>Half-life (hours)</th>
<th>Lipophilicity (log P)*</th>
<th>Protein binding (%)</th>
<th>AT_1_ receptor affinity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Losartan</td>
<td>33</td>
<td>2 (6–9)</td>
<td>–2.45</td>
<td>98.7</td>
<td>20 nmol/L</td>
</tr>
<tr>
<td>Valsartan</td>
<td>25</td>
<td>9</td>
<td>–0.95</td>
<td>95</td>
<td>1.3 nmol/L</td>
</tr>
<tr>
<td>Irbesartan</td>
<td>70</td>
<td>11–15</td>
<td>+1.48</td>
<td>90</td>
<td>2.7 nmol/L</td>
</tr>
<tr>
<td>Telmisartan</td>
<td>43</td>
<td>24</td>
<td>+1.48</td>
<td>&gt;99</td>
<td>3.7 nmol/L</td>
</tr>
<tr>
<td>Eprosartan</td>
<td>15</td>
<td>5–7</td>
<td>–</td>
<td>98</td>
<td>1.4–3.9 nmol/L</td>
</tr>
</tbody>
</table>

*The logarithm of the ratio of the concentrations of the un-ionized solute in the solvents is called log P (partition coefficient).
highest volume of distribution in its class), and high affinity for angiotensin receptor.

**PPAR-γ agonist effect of telmisartan**

Telmisartan is the only ARB which acts as a partial agonist of PPAR-γ. PPAR-γ is a ligand activated transcription factor that regulates cell growth, inflammation, lipid metabolism and insulin sensitivity. It has been successfully shown that PPAR-γ agonists limit high glucose-induced inflammation in proximal tubular cells (PTC).\(^\text{27}\) PPAR-γ agonists also exert anti-fibrotic effects in human proximal tubular cells under high glucose conditions by attenuating the increase in AP-1, TGF-β1 and downstream production of the extracellular matrix protein, fibronectin. PPAR-γ agonists can improve insulin sensitivity, reduce triglyceride levels and decrease the risk of atherosclerosis.

A study examined the ability of different ARBs to activate PPAR-γ in a cell-based transient transfection assay (Figure 2). This assay was designed to eliminate interference from endogenous nuclear receptors. In the results shown here, each ARB was used at a concentration of 10 mmol/L. Telmisartan was the only ARB to show strong (27-fold) activation, although irbesartan also showed a slight activation (2- to 3-fold). By comparison, full PPAR-γ agonists show ∼140-fold activation in this assay. Importantly, telmisartan was also the only ARB that activated PPAR-γ when tested at lower concentrations that can be achieved physiologically (1–5 mmol/L). Telmisartan therefore acts as a partial agonist of PPAR-γ.\(^\text{27}\)

![Figure 2](https://www.dovepress.com/)

**Figure 2** Inherent PPAR-γ agonist activity of various ARBs. Importantly, full PPAR-γ agonists, ie, thiazolidinediones typically have agonist activity in the 150 to 200 range. Drawn from data of.\(^\text{27}\)

There have been reports of sodium and fluid retention associated with full PPAR-γ agonist activity. Evidence suggests that PPAR-γ is highly expressed in the medullary collecting ducts with lower expression levels in glomeruli, proximal tubules and microvasculature and is a major site for increased fluid reabsorption in response to PPAR-γ agonist activity.\(^\text{41,42}\) In these collecting ducts, PPAR-γ activation increases sodium re-absorption through epithelial sodium channel (ENaC) dependent and independent mechanisms (Na⁺-K⁺-ATPase system). Guan et al showed that mice treated with thiazolidinediones (TZDs, full PPAR-γ agonists), experience early weight gain from increased total body water.\(^\text{41}\) Weight gain was blocked by the collecting duct-specific diuretic amiloride and was also prevented by deletion of PPAR-γ from the collecting duct, using PPAR-γ (flox/flox) mice. Moreover, treating cultured medullary collecting ducts with TZD increased amiloride-sensitive sodium absorption and mRNA expression of ENaC-γ isofrom through a PPAR-γ dependent pathway. Furthermore, in the blood vessels of adipose tissues, PPAR-γ ligands activate protein kinase C (PKC), vascular endothelial growth factor (VEGF), and nitric oxide (NO), which together lead to increased endothelial permeability. Thus increased renal sodium retention at the level of the collecting duct in conjunction with increased vascular permeability may cause edema development and account for the side effects of drugs with full PPAR-γ agonist activity.

The above-mentioned adverse effects have caused high attrition rates which led to the need for the discovery and characterization of alternative PPAR modulators that would retain the beneficial properties of these drugs while avoiding the cardiovascular side effects. These are called the Selective Modulators of PPAR-γ activity (SPPARγ M).\(^\text{43,46}\) PPAR-γ receptor is a multivalent receptor whose ligand binding domain can accommodate different ligands (full agonist, partial agonist, or SPPARγ M) and is capable of inducing distinct receptor combinations leading to selective gene expression. Each ligand–receptor complex assumes a somewhat different three-dimensional conformation, leading to unique and differential interactions with co-factors, histones, and other transcription factors. Consequently, each PPAR-γ ligand-receptor complex leads to a differential, but overlapping, pattern of gene expression and ultimately differential biological responses.\(^\text{43,46}\) Telmisartan has been shown to be a
partial PPAR-γ agonist and may have SPPARγMs properties as well. Schupp et al used diet-induced obese mouse models and analysis of PPAR-γ protein conformation using protease protection to show that telmisartan directly interacts with the receptor, producing distinct conformational change compared with a TZD. Telmisartan significantly reduced fasting plasma insulin and glucose levels and improved glucose tolerance and insulin sensitivity and caused significantly less weight gain compared with mice treated with pioglitazone. In vivo studies therefore suggest that Telmisartan has the beneficial properties of PPAR-γ agonists sans the side effects including sodium retention, peripheral edema and weight gain exhibited by some of the other full PPAR-γ agonists. The safety profile may imply an important use for drugs like telmisartan in the treatment of metabolic syndrome.

AMADEO trial

In the AMADEO trial, diabetic patients fulfilling the inclusion criteria were prospectively randomized (double-blind, double-dummy, forced titration) to receive telmisartan (n = 419) or losartan (n = 441), following a 2-month drug-free washout period. Patients were treated for 52 weeks in a forced-titration dosing, reaching maximum doses of 100 mg/day losartan and 80 mg/day telmisartan. Add on therapy with other antihypertensive medications were provided in case SBP remained ≥130 mmHg and DBP ≥ 80 mmHg after forced titration of the study drugs. The two study arms did not have significant differences in the use of adjunctive therapy. A drug-free follow up group included 346 patients for 2 months after the trial was concluded.

After a 4-week run-in period, 860 hypertensive patients (BP >130/80 mmHg) with type 2 diabetes and overt nephropathy (UPC ≥ 700 mg/g) were randomized to one of two treatment arms: telmisartan 40 mg or losartan 50 mg. After 2 weeks, the doses were increased to 80 mg and 100 mg, respectively. If blood-pressure control was not at goal, 25 mg of hydrochlorothiazide (HCTZ) was added, and if blood pressure was still resistant, clinicians were free to add a calcium-channel blocker. Baseline characteristics between the two treatment arms were not statistically different. After one year of therapy with the two ARBs, telmisartan provided greater reductions in the amount of protein in the urine (Table 3A), a finding not attributed to blood-pressure control, as reductions in SBP and DBP were similar in both treatment arms. As per study protocol, after a 2-month period at study completion in which both drugs were stopped, investigators reported a sustained and persistent antiproteinuric effect with the novel ARB, telmisartan. The persistent effect of telmisartan was surprising and the authors suggest that this beneficial effect could be explained by differences in receptor bindings, duration of action, or lipophilicity between the two drugs used in the trial. The difference in secondary end points of serum creatinine, eGFR, serum aldosterone and CRP failed to reach significance (Table 3B). Serum aldosterone tended to decrease to a greater extent in the telmisartan group, however statistical significance could not be attained (p = 0.07).

Does AMADEO trial unequivocally prove superiority of telmisartan?

Based on the AMADEO trial, a telmisartan-based regimen provides for robust anti-proteinuric effect in type 2 diabetic patients with hypertension and overt nephropathy. This anti-proteinuric effect is greater than a losartan-based regimen at levels of blood pressure that were not significantly different between groups and is attributed to greater lipophilicity and longer duration of action of this drug. Based on available trial data, this difference in anti-proteinuric effect may translate into better cardiovascular and renal outcomes.

The contribution of confounding factors to telmisartan’s superiority over losartan remain unclear, especially telmisartan’s partial PPAR-γ agonist effect. As discussed above, PPAR-γ agonists limit high glucose-induced inflammation in proximal tubular cells, exert anti-fibrotic effects in the same cells by attenuating the increase in AP-1, TGF-β1

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<td><strong>Secondary endpoint</strong></td>
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**Abbreviations**: CRP, C-reactive protein; eGFR, estimated glomerular filtration rate.
and the downstream production of the extracellular matrix protein fibronectin and have been shown to improve insulin sensitivity, reduce triglyceride levels and decrease the risk for atherosclerosis. This has been acknowledged by the trial authors as an explanation for telmisartan’s mechanism of action in vitro and further studies need to be done to demonstrate the action in vivo. Another issue regarding the differences in proteinuria between the groups may relate to a lower blood pressure favoring one group. The trial showed that blood pressure was decreased equally in both groups although not as much as the Captopril Study Group (CSG) (~3 mmHg in AMADEO versus ~4 mmHg in CSG). In addition, the telmisartan group separated itself from the losartan group at day 56 by 4.2 mmHg, although this difference was narrower at all other time-points. In the context that the study may be underpowered to detect differences in BP between groups, the authors concluded that the blood pressure differences were not statistically significant and larger well-designed trials were needed to detect the effects of such small blood pressure differences on proteinuria. An important observation is that both the treatment arms did not have significant side effects but there was a 20% dropout rate for the trial which was consistent in both the arms. Clarification regarding the reasons for the dropouts was not outlined in the published trial and could be important in the final analysis. Another factor that could have affected proteinuria is a disproportionate increase in dietary sodium intake in one group, since this is known to blunt the antiproteinuric effects of the ARB. However, the study detected no differences between groups in sodium excretion.

Analysis of the secondary endpoints of AMADEO trial brings forth an interesting trend. Serum aldosterone is an important part of RAAS and evidence points to some of the downstream effects of Ang II are mediated via release of aldosterone. Although not statistically significant, the study detected a trend towards greater reductions in aldosterone in the telmisartan group when compared to the losartan group (p = 0.07). The significance of this interesting trend is not clear at this point. However, future studies aimed at dissecting the role of telmisartan in ameliorating aldosterone-mediated DKD may answer this question.

![Figure 3](image_url)

**Figure 3** Primary endpoint from AMADEO trial. Patients in the two treatment groups (telmisartan vs losartan) started with the same mean UPC of 2000 mg/g creatinine. Note that the telmisartan arm had significantly lower mean UPC at every time-point over the course of the next year when compared to losartan treated patients. Drawn from data of.
Recent data from the VIVALDI trial (as described above) compared the renoprotective effects of telmisartan versus valsartan and found that they had comparable renoprotective effects. In other words, telmisartan was shown to be non-inferior at best to valsartan in slowing the progression of DKD. These findings were in contrast to the findings of the AMADEO trial. There are several possible explanations for the discrepant results including the methodology used to measure the primary endpoint; urine protein to creatinine ratio in the AMADEO trial as compared to a 24-hour urine protein used in VIVALDI trial. Furthermore, in the VIVALDI trial, participants in the valsartan group required additional antihypertensives more frequently than the telmisartan group to achieve similar levels of blood pressure control. As described above, this important confounding factor was eliminated in the AMADEO trial where none of the treatment arms was favored with better blood pressure control. Lastly, the ethnicity of the participants in the VIVALDI trial was 80% Caucasian population compared to just 45% in the AMADEO trial, the implications of this ethnic homogeneity in VIVALDI with respect to final analysis, are unclear.

**Role for aldosterone antagonism**

An important question that arises out of AMADEO trial is the use of aldosterone inhibiting agents to decrease proteinuria? Whole animal studies have suggested that aldosterone may contribute to oxidative stress, endothelial dysfunction and progressive kidney disease. Therefore, administration of selective and non-selective MR antagonists may provide additional renal benefits to the ACE inhibitors and ARBs. In a study published in 2004, Zhou et al showed that spironolactone may effectively reduce proteinuria in patients with CKD. Concerns remain with regard to the risk for hyperkalemia in patients with CKD, however. Prospective randomized trials are necessary to confirm the efficacy and safety of antagonists of aldosterone on proteinuria and progression of CKD.

**Conclusions**

Data from the AMADEO trial demonstrate the superiority and/or non-inferiority of telmisartan as a renoprotective agent when compared to losartan, an already established ARB in the treatment of DKD. The superiority as outlined is based on the unique mechanism of action of this ARB and the pharmacokinetics which seem to be extremely favorable and ideal for treating DKD. As data from recent clinical trials are emerging, use of proteinuria as a surrogate marker of renal protection is being strongly questioned. However, until additional means of measuring kidney damage on a large scale are available, this endpoint may not have been meaningfully tested. Additional questions arise due to some confounding factors in this study but the overwhelming evidence does uphold the superiority of telmisartan. Lastly, the secondary endpoint of the study does suggest a renoprotective role for aldosterone in DKD and we need well-designed randomized trials for aldosterone antagonists like spironolactone.

**Disclosures**

The authors have no conflicts of interest to disclose.

**References**