The Role of Endothelin and Endothelin Antagonists in Chronic Kidney Disease

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Keywords
Endothelin · Endothelin antagonists · Diabetic nephropathy · Hypertensive nephropathy · Focal segmental glomerulosclerosis

Abstract

Background: Endothelins (ET) are a family of peptides that act as potent vasoconstrictors and pro-fibrotic growth factors. ET-1 is integral to renal and cardiovascular pathophysiology and exerts effects via autocrine, paracrine and endocrine signaling pathways tied to regulation of aldosterone, catecholamines, and angiotensin. In the kidney, ET-1 is critical to maintaining renal perfusion and controls glomerular arteriole tone and hemodynamics. It is hypothesized that ET-1 influences the progression of chronic kidney disease (CKD), and the objective of this review is to discuss the pathophysiology, and role of ET and endothelin receptor antagonists (ERAs) in CKD. Summary: The use of ERAs in hypertensive nephropathy has the potential to decrease proteinuria, and in diabetic nephropathy has the potential to restore glycocalyx thickness, also decreasing proteinuria. Focal segmental glomerular sclerosis has no specific Food and Drug Administration-approved therapy currently, however, ERAs show promise in decreasing proteinuria and slowing tissue damage. ET-1 is a potential biomarker for autosomal dominant polycystic kidney disease progression and so it is thought that ERAs may be of some therapeutic benefit. Key Messages: Multiple studies have shown the utility of ERAs in CKD. These agents have shown to reduce blood pressure, proteinuria, and arterial stiffness. However, more clinical trials are needed, and the results of active or recently concluded studies are eagerly awaited.

Introduction

Endothelin (ET) is a 21 amino-acid peptide characterized in humans by 3 distinct genes with unique isoforms ET-1, ET-2, ET-3; all 3 are potent vasoconstrictors and pro-fibrotic growth factors \cite{1}. ET-1, however, is the predominant vascular isoform with the greatest regulatory effect on vascular tone and is integral in the regulation of renal and cardiovascular pathophysiology. ET-1 exerts effects via autocrine, paracrine and endocrine signaling pathways and regulates both catecholamines and the renin-angiotensin-aldosterone system (RAAS) \cite{2, 3}. 

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ET-1 activation is mediated by endothelin converting enzymes, which cleave ET from its precursor protein big-ET-1 into its biologically active component and C-terminal fragments [4–6]. Once activated, ET-1 has wide-ranging effects on multiple areas throughout the body, including vascular smooth muscle cells and endothelium (Fig. 1) [7]. ET-1 affects vascular smooth muscle cells following cellular uptake via clathrin-mediated endocytosis and is cleared by receptors in the lung and kidney [8–11]. ET synthesis is upregulated by gene transcription induced by angiotensin, vasopressin, interleukin-1, low extracellular pH, and cyclosporine A and can be decreased by prostacyclin, nitric oxide (NO), and natriuretic peptides [12].

ET effects are moderated by the particular isoform produced as well as the method of action utilized (paracrine, autocrine or endocrine). In addition, the downstream effect of ET is also dependent on the 2 G protein-coupled receptor isoforms (ET$_A$ and ET$_B$). In the vasculature, binding of ET$_A$ triggers vasoconstriction and mitogenesis, leading to cellular and mesangial proliferation with fibrosis. Activation of ET$_A$ also releases adrenal catecholamines and increases inotropy, elevating blood pressure (Fig. 2) [2]. ET$_B$ activation exerts an anti-hypertensive effect via vasodilation from prostaglandin, NO, and natriuresis as well as anti-mitogenic effects, which inhibit cellular proliferation and inflammation (Fig. 2) [13, 14].

In the kidney, ET-1 is a stress-induced regulator produced by the vascular endothelium, mesangial cells, and tubular epithelium with the greatest amount of activity expressed by principal cells in the medullary collecting duct [15]. ET activation can escalate progression of chronic kidney disease (CKD) via ET-1 and ET$_A$ mediated pro-fibrotic pathways, playing a role in diseases such as diabetic nephropathy (DN), hypertensive nephropathy, focal segmental glomerular sclerosis (FSGS), and autosomal dominant polycystic kidney disease (ADPKD) [16]. Early data suggest that antagonizing ET may be a promising therapeutic avenue. Preclinical and early phase clinical
studies indicate that ET antagonists may delay glomerular injury, podocyte effacement, proteinuria, and eventual scarring and sclerosis. This article will provide a comprehensive review regarding the role of ET in renal pathophysiology and specifically CKD. PubMed/Medline was searched for relevant articles using keywords “chronic kidney disease,” “CKD,” “endothelin,” “ET-1,” “endothelin antagonist,” “diabetic nephropathy,” “hypertensive nephropathy,” “focal segmental glomerular sclerosis,” and “autosomal dominant polycystic kidney disease,” and “ADPKD.” Additionally, Clinicaltrials.gov was searched for recent/current clinical trials pertaining to ET and CKD.

**ET and Renal Function**

ET has widespread effects including regulation of blood pressure, modifying cardiac output, altering systemic vascular resistance, as well as modulation of central and peripheral nervous system activity [2]. ET-1 is critical to maintaining renal perfusion and influences glomerular arteriole tone and hemodynamics; endogenous renal ET-1 is also integral to fluid and sodium homeostasis [2]. ET’s final downstream effects can be oppositional and are mediated by the receptor isoform bound. In the renal cortex, ET-1 binding of ET A drives afferent vasoconstriction decreasing renal blood flow and glomerular filtration rate (GFR). Additionally, ET-1 binding of ET A has proinflammatory and sclerotic effects on the kidney and research has shown increased transgenic expression of ET-1 in rats promotes renal scarring, increasing interstitial fibrosis and glomerulosclerosis [1, 17–20].

Alternatively, ET B receptors (ETB R) are expressed by the endothelial cells of the vasa recta and afferent arterioles where increased circulating volume-activated production of ET-1 and binding of ETB R initiates vasodilation via NO production, leading to greater perfusion of the renal medulla, decreased sodium reabsorption and natriuresis [14]. Research has shown that knockout of both ET A R and ET B R in murine collecting ducts leads to worsened hypertension as well as salt retention when compared to mice with only ET B R removed [21, 22].
There is conflicting evidence as to whether sodium transport in the proximal tubule is affected by ET [2]. However, in the medullary thick ascending limb of loop of Henle, ET-1 inhibits movement of chloride and indirectly contributes to natriuresis [7, 23, 24]. The collecting duct produces the highest levels of ET-1 in the kidney and is also the area with the highest concentration of ET receptors, with ET_B as the dominant receptor in the collecting duct and renal medulla overall [2, 25–30]. In mice, knockout of collecting duct ET-1 impaired sodium excretion, suggesting that ET-1 plays a crucial role in sodium regulation [2, 31]. This process results in diuresis via prevention of sodium and thus water reabsorption, most particularly in the collecting duct due to its high concentration of ET_B [2, 8, 9]. This experimental evidence suggests that ET-1 is necessary for routine regulation of sodium and fluid balance throughout the nephron. Loss of renal control over sodium and fluid homeostasis curtails the kidneys ability to compensate for changes in volume status and can negatively impact renal function.

**Role of ET in CKD**

Glomerular injury and scarring is the hallmark of CKD progression regardless of the underlying disease and increased production of ET-1 has been found in multiple diseases associated with CKD including diabetes and insulin resistance, obesity, and dyslipidemia [10, 31]. Renal ET-1 production is also increased by the aging process, growth factors, inflammatory cytokines, and proteinuria [31]. Schematic representation of various mechanisms by which ET-1 is involved in the causation of CKD by increased ET-1 production can be found in Figure 3. ET-1 binding of ET_A stimulates renal fibroblasts, driving increased extracellular matrix synthesis, inducing collagen production, and mesangial cell proliferation with the secretion of fibronectin and type IV collagen furthering scar formation [10, 11]. Systemic acidification stimulates renal ET-1 production, leading to acid secretion in the proximal and distal nephron as a compensatory response. Progressive renal dysfunction further worsens metabolic acidosis and ET-1 secretion contributes to CKD progression via ET-1 and ET_A mediated pro-
fibrotic pathways [16]. Furthermore, studies have demonstrated the potent vasoconstrictive effect of ET-1 on renal vascular beds. Exogenous ET-1 has resulted in decreased GFR, increased filtration fraction, as well as sodium and fluid retention [32, 33].

Recently, stress of the endoplasmic reticulum (ER) has been proposed as a mechanism of ET-1 mediated renal tubular damage. Unfolded proteins accumulate in the ER, triggering the unfolded protein response (UPR). The UPR of the ER limits transcription and translation of proteins and upregulates ER chaperone expression in an attempt to fold the proteins already collected. Over time, this pathway can lead to organ damage and cell death, as is thought to be the case in renal disease [24, 35]. Research has shown an increase in both ET-1 and ER stress in renal diseases, such as DN and acute kidney injury due to contrast, ischemia/reperfusion, or septic shock [36–44]. At this time, it is unclear whether the ET-1 upregulates the UPR or vice versa [36].

**ET and E-Receptor Antagonists in DN**

DN is one of the leading causes of CKD and end-stage renal disease (ESRD) worldwide [45]. One in 3 diabetic patients will develop DN, a disease characterized by glomerular podocyte damage with glomerular proteinuria, loss of glomerular basement membrane integrity, increased mesangial cellularity, and tubular-interstitial sclerosis [46]. Progressive glomerular damage from diabetes is augmented by ET-1 production; hyperglycemia induces ET-1, which directly degrades podocytes’ cytoskeleton leading to cellular apoptosis and podocyte effacement [47, 48]. Via upregulated ET_A receptors, the ET_A receptor/β-arrestin-1/Src-kinase complex is formed. This leads to the phosphorylation of β-catenin, increased Snail (family of zinc-finger transcription factors with a role in epithelial-mesenchymal transition [EMT]) expression and transactivation of epidermal growth factor receptor, which all contribute to podocyte damage [34, 49, 50]. Additionally, podocytes express ET receptors which when bound to ET-1 cause intracellular levels of calcium to quickly rise, influencing proliferation and signal transduction [14, 51]. ET-enhanced glomerular damage has been demonstrated in multiple animal models including the positive linear increase of ET-1 mRNA with progressive DN in rats. Well-described inflammatory mediators, including TNF-α and reactive oxygen species (ROS), have also been positively correlated with ET-1 expression and transactivation of epidermal growth factor [EMT], cell junctions are lost and collagen synthesis is diminished. EMT has been found to be essential during processes such as fibrosis, cancer development, and wound healing. Additionally, matrix metalloproteinase 9

more, ET_A and ET_B were greatly overexpressed in the renal cortex of rats with STZ induced diabetes [14].

Antagonism of ET has also improved outcomes in human studies; atrasentan, an ET_A antagonist, reduced albuminuria in patients with DN [54]. Withdrawal of atrasentan returned albuminuria to baseline. Unfortunately, due to the widespread physiological activity of ET across multiple organ systems, ET inhibition produces significant side effects and severely restricts individuals eligible for treatment and limits its clinical usefulness. For example, ET_A antagonists have been contraindicated in patients with brain natriuretic peptide > 200 pg/mL, history of heart failure, severe edema, or acute decrease in estimated GFR (eGFR) [54]. However, careful patient selection and use of diuretics may make ET inhibitors useful in CKD.

ET_A antagonists can also improve glomerular permeability through non-hemodynamic mechanisms including improved glomerular glycocalyx production [45, 55]. The glomerular glycocalyx is a glycoprotein coating on the luminal surface of glomerular capillaries that is necessary for maintaining capillary integrity. Boels et al. [55] utilized apolipoprotein E deficient mice as a model as their renovascular damage, high blood glucose, and hyperlipidemia closely replicate the phenotype of metabolic syndrome. Intraperitoneal injections of STZ were given to induce DN [45, 55]. Researchers have found that ET-1 signals increased heparanase in podocytes, an enzyme that degrades heparan sulfate glycosaminoglycans: the major constituent of glycocalyx. This led to renal damage and proteinuria, as the glomerular glycocalyx could no longer function properly [56, 57]. Mice given atrasentan had significantly reduced murine heparanase expression and demonstrated increased glomerular glycocalyx levels compared to controls [55]. In vitro studies of human umbilical vein endothelial cells showed decreased capillary glycocalyx thickness after incubation in the serum of uncontrolled diabetics. Adding atrasentan to the cell culture with diabetic human serum resulted in the restoration of glycocalyx thickness [55]. These results suggest that ET plays a critical role in the progression of DN, as antagonizing ET receptors restored glycocalyx thickness.

Tubulointerstitial fibrosis (TIF) is a common endpoint of many kidney diseases including CKD and DN. Central to TIF is epithelial cells transition into the mesenchymal phenotype, a process known as EMT. During EMT, cell junctions are lost and collagen synthesis is diminished. EMT has been found to be essential during processes such as fibrosis, cancer development, and wound healing. Additionally, matrix metalloproteinase 9
activation contributes to loss of epithelial cell junctions, disrupting the epithelial cell membrane and furthering the EMT process. Contraction of the cytoskeleton and cell motility is initiated by phosphorylation of MYPT-1, a member of the ET-1 mediating the RhoA/ROCK pathway. Lastly, blockage of phosphorylation of Yes-associated protein leads to increased transcription of mesenchymal cell markers such as αSMA. All of these aspects of EMT combined with the addition of collagen lead to tubule-interstitial fibrosis [58]. In rat studies, renal tubular epithelial cells underwent EMT in the presence of hyperglycemia; this may in part be due to the high ET-1 secretion triggered by high glucose levels [58]. Seccia et al. [58] suggest that the predominant ET receptor mediating the pathway of EMT is the ET$_B$R on proximal renal tubular cells. ET$_B$R is mostly located on endothelial cells and induces the release of NO and prostacyclin upon activation, resulting in vasodilation [59]. ET-1 binding of ET$_B$R results in decreased epithelial cell markers such as E-cadherin, while mesenchymal cell markers αSMA and vimentin increase. The role of the ET$_A$R in this process is unclear at this time [58].

**ET and Endothelin Receptor Antagonists in Hypertensive Nephropathy**

Hypertension is the second leading cause of ESRD in the United States, second only to diabetes [60]. Renal damage caused by hypertension is multifactorial and in part driven by the interplay of the RAAS and ET. Angiotensin II triggers the release of aldosterone from the adrenal cortex, which in turn increases renal ET-1 expression. Angiotensin II also directly triggers the contraction of the vascular smooth muscle of both afferent and efferent arterioles. This eventually leads to decreased renal blood flow, as well as glomerular capillary hypertension [60]. Angiotensin II also induces nicotinamide adenine dinucleotide phosphate oxidase in vascular smooth muscle cells, which drives production of ROS and increases ET-1 expression in the kidneys. ET-1 constriction of the cortex and medullary vasculature further worsens renal blood flow and promotes renal fibrosis as previously discussed. ROS production contributes to increasing arteriolar tone by decreasing NO levels and is also pro-fibrotic [60].

Currently, RAAS inhibitors are the first-line therapy for hypertensive nephropathy [60]. By inhibiting activation of angiotensin I to angiotensin II, RAAS inhibitors cut off the cytokine cascade before it amplifies resulting in improved blood pressure, decreased proteinuria, and limiting renal fibrosis [60]. Kimura et al. [61] induced hypertension in rats by giving deoxycorticosterone acetate (DOCA)-salt. The rats had their right kidney removed, and were randomly divided into 4 treatment groups: (1) DOCA-salt only, (2) nitrogen oxide synthase inhibitor Nω-nitro-L-arginine (NOARG) with DOCA-salt, (3) ET$_A$ selective receptor antagonist ABT-627, DOCA-salt and NOARG, and (4) DOCA-salt, NOARG, and nuclear factor kappa B (NF-kB) inhibitor pyrrolidinedithiocarbamate. Rats in group 2 developed the more severe renal disease compared to DOCA-salt treatment only. Nitrogen oxide synthase blockade activates NF-kB, which is genetically upstream from ET-1. Thus, upregulation of NF-kB increases ET-1 and t potentiates negative renal effects. In group 3, antagonism of ET-1 prevented all kidney damage and malfunction. In group 4, pyrrolidinedithiocarbamate usage decreased NF-kB activation, renal injury, and excess ET-1 synthesis. These results imply that the NF-kB/ET-1/ET$_A$ pathway is a major mechanism for hypertensive renal damage [61]. Additional evidence comes from a transgenic rat model of hypertension where avosentan, an ET$_A$ receptor antagonist, was beneficial in decreasing albuminuria and mortality rates compared to control rats. Furthermore, dual therapy with endothelin receptor antagonist (ERA) and valsartan (angiotensin AT1 receptor antagonist) improved outcomes compared to treatment with ERAs or valsartan alone in this rat model [62].

Despite the promising research pointing to ERAs as a possible therapeutic option in hypertensive nephropathy, the side effect profile of ERAs must be considered before clinical application. The “ASCEND” trial tested the ERA, avosentan, and was halted prior to completion due to fluid retention leading to congestive heart failure and pulmonary edema [63, 64]. Baltatu et al. [62] sought to discover whether avosentan could be useful in the treatment of hypertensive nephropathy at lower dosages in transgenic rats in hopes of limiting fluid overload. It was found that only the avosentan at the highest dose (100 mg/kg) caused the fluid overload, while the decrease in blood pressure was approximately equivalent to the change observed at the second highest dose (10 mg/kg) [62]. Rats receiving no treatment experienced a 100% mortality rate. In contrast, survival rates of transgenic rats being treated with avosentan alone as well as rats receiving both avosentan and valsartan (an angiotensin AT1 receptor antagonist) were 55.6 and 85.7% respectively [62].

**ET and ERAs in FSGS**

FSGS is a podocytopathy characterized by scarring and fibrosis of segments of glomeruli and is a significant cause of proteinuria, CKD and ESRD in the United States [65].
ET in CKD

Primary FSGS is rare in adults and accounts for 5% of ESRD cases [65]. FSGS is more common in children, but the exact incidence is not well known. Previous studies estimated a pediatric incidence of 6.9%, but more recent studies have found rates up to 23% [66]. The molecular mechanisms driving the development of FSGS are also not completely understood. Genetic mutations of structural proteins vital to podocyte architecture and function are associated with primary FSGS [67]. Alternative mechanisms include circulating factors, which damage podocytes with loss glomerular basement protein integrity and subsequent proteinuria [68]. ET has been a candidate protein and has been shown to cause pathologic changes in podocytes similar to those seen in FSGS [50]. Cultured podocytes treated with ET-1 lost cell markers indicative of podocytes including synaptopodin and acquired mesenchymal cell markers, such as aSMA. This cellular transition is similar to the EMT seen in TIF [50]. Daehn et al. [69] demonstrated that increased production of ET in FSGS also damages podocytes through oxidative stress with the injury of glomerular endothelial cells’ mitochondria and promotes adjacent podocyte apoptosis.

Despite the morbidity of FSGS, there has yet to be a specific treatment approved by the Food and Drug Administration in the United States. Therefore, the primary goal of FSGS treatment is to decrease proteinuria and first-line therapy includes a combination of corticosteroids and ACE inhibitors or angiotensin receptor blockers (ARB). Second-line therapy includes calcineurin inhibition, typically with cyclosporine or tacrolimus [70]. Antagonizing the ET system has shown promise for decreasing proteinuria and slowing renal damage in FSGS. Buelli et al. [50] utilized the Adriamycin-induced FSGS murine model and found that these mice characteristically demonstrate an elevated renal ET-1 with parietal epithelial cells activation, pseudo-crescents formation, and eventually, Adriamycin-induced injury with progression to glomerular sclerosis. Treatment with ERAs mitigated cellular injury and inflammation with significantly improved renal function in these mice.

**ET and ERAs in ADPKD**

ADPKD is a disease of autosomal dominant inheritance that consists of prolific renal cyst formation that interferes with normal renal function [71]. Incidence is estimated to range between 1:400 and 1:1,000, thus rating as one of the most commonly inherited renal disorders [72]. ET-1 is integral to cyst development and fibrotic progression of ADPKD [20, 73]. ET-1 effect on renal dysfunction in ADPKD is mediated in 3 main areas. First, ET-1 modulates gene expression in response to glomerular and tubule injury by stimulating interstitial fibroblasts to express type I collagen and smooth muscle actin. Renal cyst formation and glomerular fibrosis have thus resulted from transgenic overexpression of ET-1 in murine studies. Second, ET and its receptors are increased in ADPKD affected kidneys. In mice, ET_A mRNA is increased 5–10 times in ADPKD kidneys vs. healthy controls [74]. Research suggests that as kidney disease progresses and function decreases, urinary ET-1 is increased [32, 75]. Human ADPKD patients have increased ET-1 levels in cyst walls and fluid; furthermore, urinary levels of ET-1 are higher in ADPKD patients compared to controls [76, 77].

The association of ET with ADPKD could mechanistically be linked through hypertension, given ET’s many regulatory effects on renal and systemic blood pressure. However, Merta et al. [78] demonstrated that ET-1 and NO levels were significantly elevated in ADPKD patients vs. controls. Furthermore, there was no difference in ET-1 or NO levels when comparing normotensive to hypertensive ADPKD patients. This suggests that increased ET levels in ADPKD are independent of arterial hypertension [78]. Additional data suggested that imbalance between ET_A and ET_B receptors may be implicated in ET and disease pathogenesis of ADPKD [74]. One human study found the upregulation of ET_A receptors in renal arteries, glomeruli, and cystic epithelial cells in ADPKD patients but no significant difference in ET_B levels compared to controls. Furthermore, there is evidence that ET receptors have different effects and Chang et al. [79] demonstrated that ET_B blockade accelerated cystic disease progression while ET_A blockade leads to tubular cell proliferation.

Current treatment for ADPKD centers on symptom reduction and slowing disease progression [80]. Targeted medication includes sirolimus (mTOR inhibitor), octreotide (a somatostatin analog), and/or tolvaptan (a selective vasopressin V2 receptor antagonist) [81]. Sirolimus and similar drug everolimus have been shown to decrease the rate at which total kidney volume (TKV) increases, however current evidence suggests they do not slow disease progression and renal damage as measured by eGFR [81–83]. Octreotide also slows increasing TKV, but like sirolimus, it has no effect on decreasing renal function [81, 84–86]. Tolvaptan has also been shown to slow increasing TKV and Torres et al. [87] demonstrated a significant reduction in loss of kidney function vs. placebo, with declining kidney function measured as a 25% decrease in the reciprocal serum creatinine level [81].

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Kidney Dis

ET in CKD

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ET-1 also has utility as a noninvasive biomarker to identify disease progression in high-risk ADPKD patients. This is especially useful since disease markers such as eGFR do not change until significant renal damage has already occurred [77, 88]. Raina et al. [77] found that urinary ET-1 positively correlated with TKV, which serves as a marker for cyst enlargement (r = 0.426), however, this correlation was not statistically significant (p = 0.1) and warrants further clinical investigation.

Clinical Trials Evaluating ERAs for CKD Management

All the available clinical trials involving ERAs for CKD management are summarized in Table 1. In a study by Kohan et al. [89], it was found that atrasentan at doses of 0.75 mg and 1.75 mg/day significantly improved residual albuminuria in type II diabetes mellitus patients who were already on renin-angiotensin system blockers. Short-term results of the ASCEND study investigating the effect of ERA, avosentan, on progression of overt DN showed that avosentan significantly reduced albuminuria when added to renin-angiotensin system blockers. However, the trial was terminated prematurely after a median follow-up of 4 months due to higher cardiovascular adverse events, including fluid overload, congestive heart failure, and death [64].

The “Study of DN with Atrasentan” (SONAR) phase III trial was meant to determine the efficacy of atrasentan (an ERA) to treat type II DN and delay progression of kidney disease by decreasing the urine albumin-to-creatinine ratio (UACR) [90]. This trial was different than ASCEND and similar studies as it utilized a response enrichment design, where the trial only included patients likely to benefit from the experiment. Subjects included in the trial went through a Run-In Period to optimize RAS inhibitor dosage followed with the enrichment period with atrasentan to determine UACR. Afterwards, responders and non-responders were included in a double-blind treatment period [91, 92]. However, AbbVie terminated this study in 2018 due to lower than expected renal end points; not due to any safety concerns for the participants [93]. Another study was the “Zibotentan Better Renal Scleroderma Outcome Study”, which ran from October 2014 to 2017 per ClinicalTrials.gov. Zibotentan Better Renal Scleroderma Outcome Study was a phase II investigation of zibotentan, an ERA, and its effect on the progression of kidney disease due to scleroderma. The primary outcome measurement was soluble vascular cell adhesion molecule 1 level, which are reflective of renal involvement in scleroderma. Whether these levels were measured via the blood or the urine is unclear and the results of the study have not yet been published [99]. A different study by Bérèzne et al. [100] tested the effect of bosentan in scleroderma renal crisis patients and found no improvement in renal outcomes.

Side Effects of ERAs

The side effect profile of ERAs is substantial and may limit their clinical usefulness. The most common adverse effect among ET blockers is fluid retention, which is due to ET\textsubscript{B}/ET\textsubscript{A} blockade. This blockade leads to a decrease in NO and PGE\textsubscript{2} causing an increase in sodium and water reabsorption. “Dual” antagonists have an ET\textsubscript{A}: ET\textsubscript{B} selectivity <100-fold while selective antagonists work on ET\textsubscript{A}, thus
Table 1. Clinical trials of endothelin antagonists in chronic kidney disease conditions

<table>
<thead>
<tr>
<th>Study</th>
<th>Clinicaltrials.gov registration no.</th>
<th>Number of subjects (n) and age</th>
<th>Study type/phase</th>
<th>Conditions</th>
<th>Interventions</th>
<th>Study status</th>
<th>Conclusions/outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zibotentan Better Renal Scleroderma Outcome Study (ZEBRA) [99]</td>
<td>NCT02047708</td>
<td>( n = 27 ) Eligibility ≥18 years</td>
<td>Phase II RCT</td>
<td>Scleroderma, scleroderma renal crisis, CKD</td>
<td>Zibotentan (ZD4054)</td>
<td>Completed in October 2017</td>
<td>Results not found/published yet</td>
</tr>
<tr>
<td>Study of Diabetic Nephropathy with Atrasentan (SONAR) [90]</td>
<td>NCT01858532</td>
<td>( n = 5,112 ) Eligibility 18–85 years</td>
<td>Phase III RCT</td>
<td>Diabetic nephropathy</td>
<td>Atrasentan, placebo</td>
<td>Terminated</td>
<td>Study was terminated due to lack of promising evidence, not due to safety concerns, per report from pharmaceutical company</td>
</tr>
<tr>
<td>Trachtman et al. [97], 2018 (DUET study)</td>
<td>NCT01613118</td>
<td>( n = 109 ) 8–75 years</td>
<td>Phase II RCT</td>
<td>Focal segmental glomerulosclerosis</td>
<td>Sparsentan, irbesartan</td>
<td>Completed</td>
<td>Sparsentan was safe, well tolerated and significantly reduced proteinuria compared to irbesartan</td>
</tr>
<tr>
<td>Lin et al. [95], 2018</td>
<td>NCT01424319; NCT01356849; NCT0139580</td>
<td>( n = 257 ) 64–67 years (mean age)</td>
<td>Results from 3 phase II RCTs</td>
<td>CKD, diabetic nephropathy</td>
<td>Atrasentan, placebo</td>
<td>Completed</td>
<td>No significant difference in incidence of peripheral edema between placebo and atrasentan treated groups; Further, results suggested that a maximum decrease of 46% in urine albumin creatinine ratio would be achieved</td>
</tr>
<tr>
<td>Webb et al. [96], 2017</td>
<td>NCT01399580</td>
<td>( n = 48 ) 53–77 years</td>
<td>Phase IIb RCT</td>
<td>CKD, diabetic nephropathy</td>
<td>Atrasentan, placebo</td>
<td>Completed</td>
<td>Found that thoracic bioimpedance did not correlate with edema or increased weight</td>
</tr>
<tr>
<td>Bérézne et al. [100], 2017 (abstract)</td>
<td>NCT01241383</td>
<td>( n = 16 ) 46.3–71.3 years</td>
<td>Phase II open label</td>
<td>Scleroderma renal crisis</td>
<td>Bosentan</td>
<td>Completed</td>
<td>Bosentan did not improve renal outcomes</td>
</tr>
<tr>
<td>Kohan et al. [89], 2011</td>
<td>NCT00920764</td>
<td>( n = 89 ) 51–77 years</td>
<td>Phase IIa RCT</td>
<td>CKD, diabetic nephropathy</td>
<td>Placebo, atrasentan</td>
<td>Completed</td>
<td>Atrasentan is safe and effective in decreasing residual albuminuria significantly in patients with type II diabetes mellitus already on renin angiotensin system inhibitors</td>
</tr>
<tr>
<td>Dhaun et al. [98], 2011</td>
<td>NCT00810732</td>
<td>( n = 27 ) 36–60 years</td>
<td>Phase II RCT</td>
<td>Nondiabetic CKD with proteinuria (&gt;300 mg/day)</td>
<td>Sitaxsentan, nifedipine, placebo</td>
<td>Completed</td>
<td>Sitaxsentan significantly decreased 24-h proteinuria, protein:creatinine ratio, blood pressure, and pulse wave velocity compared to placebo; Nifedipine did not reduce proteinuria</td>
</tr>
<tr>
<td>Endothelin Receptor Antagonism in Proteinuric Nephropathy</td>
<td>NCT00722215</td>
<td>( n = 22 ) Eligibility 18–70 years</td>
<td>Phase I RCT</td>
<td>CKD with proteinuria</td>
<td>Selective endothelin A receptor antagonist (BQ-123), 0.9% saline, nifedipine</td>
<td>Completed</td>
<td>Results not found</td>
</tr>
<tr>
<td>Bosentan Use in Patients with Diabetic Nephropathy [94]</td>
<td>NCT00638131</td>
<td>( n = 1 ) Eligibility 18–80 years</td>
<td>Phase III RCT</td>
<td>Type 2 diabetes</td>
<td>Bosentan</td>
<td>Terminated</td>
<td>Results not found; Reason for termination unclear</td>
</tr>
<tr>
<td>Mann et al. [64], 2010 (ASCEND study)</td>
<td>NCT00120328</td>
<td>( n = 1392 ) 50–70 years</td>
<td>Phase III RCT</td>
<td>Diabetic nephropathy</td>
<td>Avoosentan</td>
<td>Terminated</td>
<td>Short-term results showed that avosentan significantly reduced albuminuria in type II diabetes or overt nephropathy patients who were already on renin angiotensin system blockers; However, trial was terminated after median follow-up of 4 months due to higher cardiovascular adverse events, including fluid overload, congestive heart failure and death</td>
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</tbody>
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theoretically resulting in less fluid retention. Individual ERAs have specific side effects that must be monitored including hepatotoxicity with patients on sulfonamide-based ERAs [101–103]. ET<sub>A</sub>/ET<sub>B</sub> antagonism is teratogenic, thus contraindicated during pregnancy, but this may be less of a concern in the aging CKD population [21]. Testicular toxicity is another rare, but serious side effect [31]. There are strict patient populations to be avoided due to the many side effects of ERAs including severe CKD and congestive heart failure which may complicate fluid retention management [31]. The “ASCEND” trial tested avosentan but was halted due to the dangerous retention of fluids [63, 64]. SONAR trial showed good outcomes when ERAs were used with diuretics [90]. Thus, a therapeutic potential may be possible at either lower dosages or if ERA’s were utilized in concert with CKD fluid management strategies aimed at ameliorating the dangerous side effects found in this drug class. In addition, any future clinical trials of ERAs for CKD will likely need to include ERA in addition to rather than in place of standard of care, which at present includes an ACEi/ARB, but could potentially be expanded to SGLT2i in the near future.

**Conclusion**

ETs are a family of potent vasoconstrictors and profibrotic growth factors and its receptors (ET<sub>A</sub> and ET<sub>B</sub>) are integral in the regulation of renal and cardiovascular pathophysiology. However, ET activation can escalate the progression of CKD and data has suggested that antagonizing ET may be a promising therapeutic avenue. Multiple studies have shown the utility of ERAs in both diabetic and nondiabetic CKD by reducing blood pressure, proteinuria, and arterial stiffness. However, there have been no studies to date demonstrating that ERAs improve clinical outcomes in CKD. Multiple large clinical trials are needed, and the results of active or recently concluded studies are eagerly awaited for analysis and comparison to current data.

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