Urgent call for reconsideration of chronic kidney disease

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Abstract
Circulating toxins namely: free radicals, cytokines and metabolic products induce glomerular endothelial dysfunction, hemodynamic maladjustment and chronic ischemic state; this leads to tubulointerstitial fibrosis in chronic kidney disease (CKD). Altered vascular homeostasis observed in late stage CKD revealed defective angiogenesis and impaired nitric oxide production explaining therapeutic resistance to vasodilator treatment in late stage CKD. Under current practice, CKD patients are diagnosed and treated at a rather late stage due to the lack of sensitivity of the diagnostic markers available. This suggests the need for an alternative therapeutic strategy implementing the therapeutic approach at an early stage. This view is supported by the normal or mildly impaired vascular homeostasis observed in early stage CKD. Treatment at this early stage can potentially enhance renal perfusion, correct the renal ischemic state and restore renal function. Thus, this alternative therapeutic approach would effectively prevent end-stage renal disease.

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Key words: Hemodynamics; Vascular homeostasis; Early diagnostic markers; Chronic kidney disease; Vasodilators

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INTRODUCTION
Homeostasis of the vital organs in the body depends on the balance between nutrient supply through vascular perfusion and the integrity of structure and function of the organs. In this regard, kidney integrity depends mainly on the renal vascular supply. Philosophically, normal homeostasis of the kidney follows the so-called Natural Wisdom “The Middle Tract is the Balance of Nature”, which implies that the normal integrity of the kidney depends on normal vascular perfusion. Any deviation of blood perfusion, either too much or too little would be harmful to the kidney[1]. Such wisdom can be illustrated by the correlation between blood perfusion and its organ’s structure and function. Under normal circumstances, the intact tubulointerstitium has been shown to be surrounded by an adequate supply of peritubular capillary plexus. In contrast, under pathological conditions, such as in chronic kidney disease (CKD), the normal tubulointerstitial structure is replaced by tubulointerstitial fibrosis, along with the disappearance of peritubular capillary plexus which is replaced by renal microvascular disease[2,3]. The spatial relationship between renal perfusion and kidney integrity will be the context of the following issues: (1) Renal microvascular disease and tubulointerstitial fibrosis; (2) Why does the present therapeutic strategy fail to restore renal function in CKD? (a) CKD is recognized and treated at a
rather late stage; and (b) Altered vascular homeostasis in late CKD; and (3) An innovative therapeutic strategy to implement the treatment at early stage CKD.

**RENA L MICROVASCULAR INJURY AND TUBULOINTERSTITIAL FIBROSIS**

Accumulating evidence supports the suggestion that there are toxins such as free radicals, cytokines and metabolic products circulating through the renal microcirculation in a variety of CKDs. Abnormally elevated oxidant and antioxidant deficiencies have been repeatedly documented in both mild as well as severe forms of CKD.  

The circulating toxins can induce injury to renal microvasculature. Such vascular injury detaches the endothelial cell from the vascular wall and this is reflected in an increase in the number of circulating endothelial cells in a variety of CKD patients. In addition to the increased number of circulating endothelial cells, the remaining endothelial cells also become dysfunctional. Glomerular endothelial dysfunction is characterized by upregulation of vasoconstrictors such as angiotensin II, endothelin, thromboxane A2, adhesion molecules, procoagulant activity and reactive oxygen species. Enhanced expression of vasoconstrictors induces hemodynamic maladjustment characterized by a preferential constriction at the efferent arteriole and thus a corresponding reduction in peritubular capillary flow supplying the tubulointerstitial structure (Figure 1). This phenomenon is well documented in CKD. It is interesting to observe that an intact tubulointerstitial structure is usually associated with normal level of peritubular capillary flow. Reduced peritubular capillary flow has been observed in all CKD patients. A mild reduction in peritubular capillary flow has been noted to precede the development of tubulointerstitial fibrosis. An increased reduction in peritubular capillary flow leads to the appearance of tubulointerstitial fibrosis. A further reduction in peritubular capillary flow is associated with a higher degree of tubulointerstitial fibrosis (Table 1). This suggests that the reduction in peritubular capillary flow determines the development of tubulointerstitial fibrosis. In addition, the dysfunctioning endothelial cell expresses procoagulant activity which is reflected in blood hypercoagulability, blood hyperviscosity, a shortened platelet half life and a shortened fibrinogen half life, indicating an increased consumption of local intravascular coagulation, plausibly in the renal microcirculation. Thus correcting both the altered blood coagulability with anticoagulant and antiplatelet agents, as well as correcting the chronic ischemic state with vasodilators is an appropriate therapeutic target.

**WHY DOES THE PRESENT THERAPEUTIC STRATEGY FAIL TO RESTORE RENAL FUNCTION IN CKD?**

**Table 1** A correlation between peritubular capillary flow reduction and tubulointerstitial fibrosis

| Clinical setting     | Peritubular capillary flow mL/min per 1.73 m² | Tubulointerstitial fibrosis |
|----------------------|---------------------------------------------|----------------------------|
| Normal               | 480                                         | Negative                   |
| Early CKD            | 250-400                                     | +                           |
| Late CKD             | < 250                                       | ++++                       |

CKD: Chronic kidney disease.

[Figure 1] Hemodynamic maladjustment induces chronic renal ischemia and tubulointerstitial fibrosis.

Only patients associated with creatinine clearance under 60 mL/min per 1.73 m², or serum creatinine greater than 1 mg/dL, recognition of CKD is practically limited to late CKD (stages 3-5) since serum creatinine does not change until the creatinine clearance drops to the 50% level. This implies that treatment of CKD is usually initiated at a rather late stage. Early stage CKD patients have generally been left untreated, and the disease allowed to progress without any appropriate therapeutic intervention. Treatment of these CKD patients with vasodilators shows therapeutic unresponsiveness and fails to correct the chronic ischemic state. This issue leads us to propose that vascular homeostasis explains such therapeutic failure.

**Altered vascular homeostasis in late CKD**

It has been recently demonstrated that altered vascular homeostasis and impaired nitric oxide (NO) production are responsible for therapeutic resistance to vasodilators in late CKD. With respect to vascular homeostasis, a normal vascular homeostasis is the balance between vascular injury and vascular repair (Figure 2). Under normal circumstances, vascular injury results in an increased number of endothelial cells detaching from the diseased vascular wall into the circulation, so-called circulating endothelial cells which express receptor-bound vascular endothelial growth factor (VEGF) as suggested by Hohenstein et al. Such vascular injury would trigger vascular repair by recruiting angiogenic factors such as VEGF which would activate through VEGF receptor 1 (VEGFR 1) inducing Akt phosphorylation, coupled with endothelial nitric oxide synthase (eNOS), and enhanced NO production. Enhanced
Enhanced antiangiogenesis would induce a thickening of the vascular wall, oxidative stress, NFKB and p38, JAK STAT and eventually endothelial cell dysfunction, activating NADPH oxidase, generated by the upregulation of angiotensin II. With respect to defective angiogenic factors, as indicated on the left hand side of Figure 3, the defective VEGF and VEGFR 1 would impair the Akt phosphorylation, uncouple eNOS, and impair NO production. An impaired NO production in conjunction with defective endothelial progenitor cells and defective angiopoietin 1 would impair the physiological stimulation of endothelial cell proliferation and maturation. They would integrate together in an insufficient vasculogenesis and vascular repair. With respect to the abnormally elevated angiogenic factors, VEGF activates through VEGFR 2 inducing abnormal Akt phosphorylation by an NO-independent pathway, resulting in a proliferation of abnormally immature endothelial cells. The presence of defective angiopoietin 1, is responsible for the immature endothelial cell proliferation. In addition, the presence of abnormally elevated angiopoietin 2, endothelial cells would be further destabilized and endothelial apoptosis induced. Collectively, they would integrate in the formation of abnormally immature endothelial cells, which would be consistent with endothelial myofibroblast transition cell indicated by Li et al. With respect to the vascular smooth muscle cell (VSMC) proliferation, this is triggered by the upregulation of angiotensin II secondary to endothelial cell dysfunction, activating NADPH oxidase, oxidative stress, NFKB and p38, JAK STAT and eventually stimulating VSMC proliferation.

The alternative therapeutic strategy would focus on early stage CKD patients, who mostly have been untreated or received inappropriate treatment. Our recent study on vascular homeostasis in early stage CKD supports this concept. In type 2 diabetic nephropathy, the vascular homeostasis observed in the normoalbuminuric stage indicated that both angiogenic factors namely: VEGF, angiopoietin 1 and VEGF receptor 1; as well as antiangiogenic factors namely: angiopoietin 2 and VEGF receptor 2, were within normal limits. In non-diabetic early stage CKD patients, the vascular homeostasis indicated that angiopoietin 1 was the only angiogenic factor showing a mild decrease, and that angiopoietin 2 was the only antiangiogenic factor showing a mild elevation. Thus these findings render support the theory that vascular narrowing of the vascular lumen, and eventually a reduction in vascular perfusion, which eventually leading to the development of neoangiogenesis and progressive vascular disease. The altered vascular homeostasis observed in late stage CKD indicates both an insufficient vasculogenesis associated with an impaired NO production, which explains the therapeutic resistance to vasodilators, as well as the clinical progression of renal microvascular disease of increasing severity. Also, the progression of renal microvascular disease correlates with the altered renal hemodynamics characterized by a progressive reduction in peritubular capillary flow along with a progressive decline in renal function. Such therapeutic failure of current practice with vasodilators in late stage CKD requires an alternative strategy to focus the treatment at the new target group of CKD patients at the early stage of renal function impairment (Figure 3).

**AN ALTERNATIVE THERAPEUTIC STRATEGY TO IMPLEMENT TREATMENT AT AN EARLY CKD STAGE**

The alternative therapeutic strategy would focus on early stage CKD patients who have been untreated or received inappropriate treatment. Our recent study on vascular homeostasis in early stage CKD supports this concept. In type 2 diabetic nephropathy, the vascular homeostasis observed in the normoalbuminuric stage indicated that both angiogenic factors namely: VEGF, angiopoietin 1 and VEGF receptor 1; as well as antiangiogenic factors namely: angiopoietin 2 and VEGF receptor 2, were within normal limits. In non-diabetic early stage CKD patients, the vascular homeostasis indicated that angiopoietin 1 was the only angiogenic factor showing a mild decrease, and that angiopoietin 2 was the only antiangiogenic factor showing a mild elevation. Thus these findings render support the theory that vascular
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homeostasis would likely be adequately functional in early stage CKD. With the adequate vasculogenesis observed in early stage CKD, vasodilator treatment would relax the efferent arteriole by enhancing the peritubular capillary flow. Increased peritubular capillary flow would inhibit the process of tubulointerstitial fibrosis indicated by the decline in Fe-Mg value following vasodilator treatment - an index indicating renal regeneration. Vasodilator treatment would also relax the afferent arteriole and thereby increase the glomerular filtration rate - an index indicating renal function improvement. In fact, therapeutic implementation with appropriate vasodilators at this early stage in CKD has indeed been able to enhance peritubular capillary flow, as well restore renal function.

In conclusion, therapeutic implementation of vasodilator treatment in early stage CKD, in an environment favorable to renal angiogenesis and regeneration, can effectively prevent end-stage renal disease.

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