Brief Review

Cellular and molecular aspects of diabetic nephropathy; the role of VEGF-A

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\textbf{A B S T R A C T}

The prevalence of diabetes mellitus increased during the last century and it is estimated that 45% of the patients are not diagnosed. In South America the prevalence of diabetes and chronic kidney disease (CKD) increased, with a great disparity among the countries with respect to access to dialysis. In Ecuador it is one of the main causes of mortality, principally in the provinces located on the coast of the Pacific Ocean. The greatest single cause of beginning dialysis is diabetic nephropathy (DN). Even using the best therapeutic options for DN, the residual risk of proteinuria and of terminal CKD remains high. In this review we indicate the importance of the problem globally and in our region. We analyse relevant cellular and molecular studies that illustrate the crucial significance of glomerular events in DN development and evolution and in insulin resistance. We include basic anatomical, pathophysiological and clinical concepts, with special attention to the role of angiogenic factors such as the vascular endothelial growth factor (VEGF-A) and their relationship to the insulin receptor, endothelial isoform of nitric oxide synthase (eNOS) and angiopoietins. We also propose various pathways that have therapeutic potential in our opinion. Greater in-depth study of VEGF-A and angiopoietins, the state of glomerular VEGF resistance, the relationship of VEGF receptor 2/nephrin, VEGF/insulin receptors/nephrin and the relationship of VEGF/eNOS-NO at glomerular level could provide solutions to the pressing world problem of DN and generate new treatment alternatives.

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La prevalencia de diabetes mellitus aumentó en el último siglo y se estima que el 45% de los pacientes, no estarían diagnosticados. En Sudamérica la prevalencia de diabetes y de enfermedad renal crónica (ERC) incrementó, existiendo gran disparidad entre los países respecto al acceso a diálisis. En Ecuador es una de las principales causas de mortalidad, principalmente en las provincias ubicadas en la costa del océano Pacífico. La mayor causa aislada de ingreso a diálisis es la nefropatía diabética (ND). Aun utilizando las mejores opciones terapéuticas para la ND, el riesgo residual de proteinuria y de ERC terminal permanece elevado. En esta revisión describimos la importancia del problema en el mundo y en nuestra región. Analizamos estudios moleculares y celulares relevantes que indican la crucial importancia de eventos glomerulares en el desarrollo y en la evolución de la ND y en la insulinoresistencia. Incluimos conceptos anatómicos, fisiopatológicos y clínicos básicos, desarrollando especial énfasis en el rol de factores angiogénicos como el factor de crecimiento vascular endotelial (VEGF-A) y su relación con el receptor de insulina, la sintasa endotelial de óxido nítrico-óxido nítrico (eNOS) y las angiopoietinas. En el transcurso del texto proponemos diversas vías, que a nuestro entender tienen potencial terapéutico. Profundizar en el estudio del VEGF-A y las angiopoietinas, el estado de VEGF resistencia glomerular, la relación del receptor 2 de VEGF/nefrina, VEGF/receptores de insulina/nefrina, la relación VEGF/eNOS-ON a nivel glomerular podría aportar soluciones al acuciante problema de la ND en el mundo y generar nuevas alternativas de tratamiento.

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the insulin receptor and angiopoietins. Finally, we will consider basic aspects and the analyses of recently published molecular and cellular studies.

**Anatomical and pathophysiological aspects of DN**

Diabetes involves functional and structural kidney alterations that induce proteinuria at variable magnitudes, ranging from micrograms to several grams per day\(^7,8\). The risk of developing ECKD is related to albumin urinary excretion, and early treatment with RAAS inhibitors is important due to the beneficial renal and systemic effects\(^7,8\). DN is accompanied by persistent albumin urinary excretion or microalbuminuria, which is defined as the loss of urinary albumin ranging from 20 to 199 µg/min or 30 to 299 mg/d on two different occasions and when the albumin/creatinine ratio is 30-299 mg/g in an isolated urine sample\(^7,8\). In type 1 diabetes, albumin urinary excretion should be quantified on an annual basis, from the fifth year following diagnosis onwards; in type 2 diabetes, given the difficulty to accurately state its onset, measurement is preferable from the moment the disease is diagnosed\(^7,8\). In one study, the prevalence of microalbuminuria in patients with type 2 diabetes was 24.9% after a ten-year follow-up\(^4\), but 30% of patients with type 2 diabetes and no microalbuminuria developed DN. It is also important to quantify glomerular filtration (GF), since some patients only show renal function impairment with no signs of proteinuria\(^7,8\).

Considering that 85% of the patients with diabetes have type 2 diabetes, better biomarkers are required\(^7,8\). Risk factors contributing to the development of DN are hyperglycaemia, hypertension (HTN), dyslipidaemia, age over 65 years, male gender, smoking habit, family history and Hispanic or Afro-American origin\(^7,8\). Familial clustering was reported in populations with different ancestors, especially in Pima Indians and Afro-Americans\(^6\). Mooyaart et al. found 24 genetic variations associated with DN\(^7\). Epigenetic mechanisms were also implied\(^6,13\). For example, chronic hyperglycaemia, without altering the nucleotide sequence, may modify DNA or methylate histones associated with DNA\(^18\). However, the significance of these findings on the development of DN has not been determined yet.

Many factors were implied in DN pathophysiology, such as: glucose, glucose receptors, VEGF-A, NO, reactive oxygen species (ROS), transforming growth factor beta (TGF-Beta), RAAS, kinin-kallikrein system, mammalian target of rapamycin, inflammation, tumour necrosis factor alpha, adiponectin, advanced glycation end products and receptors thereof, mitochondrial oxidative stress and microRNA\(^18\).

From the pathologic point of view, type 1 and type 2 diabetes induce common kidney lesions. These lesions were characterised in type 1 diabetes\(^7,8\). In type 2 diabetes, the kidney histology and course have special features, associated with comorbidities such as HTN, vascular diseases, ageing and obesity\(^7,8\). Five years after diabetes diagnosis, there is hyperfiltration, microalbuminuria, glomerulomegaly, glomerular basal membrane (GBM) thickening and alteration of podocytes\(^26\). Subsequently, the extracellular matrix (ECM) is deposited in the mesangium. Approximately ten years later, proteinuria and HTN are evident, and GF becomes progressively impaired\(^7,23,24\). Within a period of 20 to 25 years, sclerosis is advanced, there is tubulointerstitial fibrosis and CKD progresses to end-stage phases\(^7,24,26\).

Meanwhile, the glomeruli, tubules, interstitium and renal arteries are modified by the diabetic environment. Glomerular changes involve the glomerular filtration barrier (GBF), ECM, and the main cells composing it (podocytes, endothelial cells and mesangial cells)\(^7,16,19-21\). In addition, it prevents the abnormal passage of plasma protein based on size and load, and its alteration was associated with proteinuria\(^7,15,19,20,25\). The GBF is composed of podocytes, GBM, and the endothelium (Figure 2). Podocytes are markedly differentiated epithelial cells, with a large cell body, and primary and secondary extensions connected by slit diaphragms (SD)\(^15,19,20\). The SD is permeable to water and small solutes, but it is selective to large molecule passage, which is a key factor in GBF permeability\(^25\). Moreover, it is composed of a protein complex, where nephrin plays an important role\(^7,15,19,20\). On the apical side, podocytes float within the urinary space, while on the basolateral side, they make contact with the GBM. Podocyte cytoskeleton proteins are related to GBM proteins through integrins and dystroglycans\(^15,20,25\). The GBM is mainly composed of proteins, such as collagen IV and laminins\(^15,25\). The fenestrated endothelium, covered by glyocalyx, is the inner most layer of the GBF\(^2,15,21,25\). Diabetes alters the three layers that make up the GBF. Among the early changes, neoangiogenesis in the glomerular vascular pole and loss of endothelial fenestrations have been described\(^7,16,22,23\). The GBM shows an increased thickness due to protein exchange alterations\(^7,15,19,20,25\).

![Mortality due to Diabetes Mellitus](image)

Figure 1 – In Ecuador, mortality caused by diabetes mellitus was higher in the provinces of Guayas, Los Ríos and Manabí, located on the Pacific coast. Map shows the mortality rate due to diabetes mellitus (deaths/100,000 individuals per year, INEC [Instituto Nacional de Estadísticas y Censos - National Institute of Statistics and Census of Ecuador] 2011). This figure is part of a figure originally published by Neira-Mosquera et al.\(^6\), with minor modifications (authorised reproduction).
VEGF-A is a potent angiogenic factor related to normal and pathological angiogenesis. It promotes the proliferation, differentiation and migration of endothelial cells; it induces vasodilation and increases vascular permeability. It plays an important role in kidney development; in the adult kidney, it is secreted by podocytes and is essential for the maintenance of the GFB. It acts through tyrosine-kinase receptors, which are known as VEGF receptor 1 and 2 (VEGFR1 and 2). VEGFR2 is expressed in endothelial cells and podocytes; it is related to the most important signals of VEGF-A. Two co-receptors called neuropilins 1 and 2 amplify the VEGFR2 signal.

There is evidence that glucose directly and indirectly stimulates VEGF-A expression in podocytes through angiotensin II and TGF-Beta. Glucose plays a very important role in DN pathophysiogenesis. Glycaemic control reduces DN progression and induces reversion of proteinuria and advanced histological lesions. In a 30-year follow-up study, proteinuria and nephropathy were reduced by the strict control of glycaemia. The role of VEGF-A in DN

Figure 2 – Schematic representation of the glomerulus, the glomerular filtration barrier (GFB) composed of podocytes (P), the glomerular basal membrane (GBM), and the endothelium. Plasma ultrafiltration passes through the GFB (black arrow) to reach the urinary space (US). Podocytes (green) make contact with several glomerular capillaries (represented as red circles) and the intraglomerular mesangium (M). The GBM (black line) wraps the capillaries and surrounds the mesangium. The glomerular endothelium is represented by a discontinuous light-blue line, located between the capillary lumen (CL) and the GBM, the vascular pole in the lower part of the glomerulus, the tubular pole in the upper part. B: Ultrastructure of the GFB observed with an electronic microscope: podocytes, GBM, slit diaphragm (SD) and fenestrated endothelium. Plasma ultrafiltration passes through the GFB (yellow arrow) from the capillary lumen (CL) towards the urinary space (US).
uria, GF and HTN showed an improvement in patients with type 1 diabetes when there was better glycaemic control. With a higher control of hyperglycaemia, GBM has shown less thickening. Histological changes of advanced DN reverted 10 years after pancreas transplantation. Haraguchi et al. were able to revert nephritic-range proteinuria and histological lesions compatible with advanced DN after five years of intensive treatment of hyperglycaemia. Treatment with bariatric surgeries administered to patients with type 2 diabetes and obesity improved GF and proteinuria, which was related to weight loss and decreased hyperglycaemia.

Hyperglycaemia increases renin and angiotensinogen expression in mesangial cells. Mesangial cells and podocytes synthesise angiotensin II and express angiotensin receptors. The increase in angiotensin II stimulates the expression of TGF-Beta, VEGF-A, connective tissue growth factor (CTGF), interleukin 6 and chemoattractant protein for monocytes-1 inducing expansion of the ECM and podocyte apoptosis. CTGF; the increase in VEGF-A inhibits TGF-Beta expression, in a negative feedback mechanism. In contrast, the increase in VEGF-A in diabetes is associated with elevated TGF-Beta and CTGF, proliferation and build-up of proteins in the glomerular ECM. TGF-Beta has been related to the proliferation of mesangial cells, diffuse nodular glomerulosclerosis and also fibrosis. In transgenic mice with no TGF-Beta type 2 receptor and the administration of anti-TGF-Beta antibodies prevented mesangial build-up and kidney function impairment. These antibodies represent a therapeutic hope for DN, but they are not available for human use yet.

**Glomerular VEGF-A modifications in DN**

Starting from early DN stages, systemic and renal VEGF-A are elevated in humans and mice, VEGF-A has been associated with neoangigenesis. RAAS, VEGF-A and nephrinuria were seen to be involved in this process. Cultured podocytes and endothelial cells increased VEGF-A and VEGFR2 expression in response to the increase in glucose. We showed that glomerular VEGF is a key factor for DN development and progress. Normoglycaemic mice with VEGF overexpression in podocytes developed glomerulomegaly, hyperfiltration, GBM thickening and podocyte lesion, which are changes similar to early DN. In these transgenic mice, diabetes caused massive proteinuria, advanced nodular glomerulosclerosis and less nephrin expression. Diabetic mice with no VEGF overexpression only showed mild diffuse glomerulosclerosis. These experiments demonstrate that the increase in glomerular VEGF, irrespective of the diabetic environment, generates identical changes to the early DN and that increasing glomerular VEGF speeds up DN progress to more advanced stages. In the absence of diabetes, the urinary VEGF-A was reported to be a good marker of glomerular expression or DN severity. Urine and systemic VEGF-A levels were high in diabetic mice with and without glomerular VEGF overexpression. Probably, within the diabetes context, urinary excretion of VEGF-A reflects systemic levels, while hiding VEGF glomerular changes. In short, these experiments suggest that glomerular VEGF-A is a determining factor in DN, that VEGF overexpression in podocytes is dangerous, and that glucose directly and indirectly stimulates the VEGF-A signalling cascade in podocytes. In diabetes, urinary and systemic VEGF-A did not correlate with either glomerular VEGF expression or with the severity of glomerular lesions, which brings into question the use of VEGF-A as a DN biomarker.

**Glomerular VEGF-A reduction was shown to generate GF lesions, proteinuria and kidney failure in animals and humans.** Transgenic mice with silencing of VEGF-A in podocytes showed AKI, alteration of the three GBF layers and reduced integrin expression. Some patients treated with anti-VEGF-A antibodies showed proteinuria, endothelial lesions and thrombotic microangiopathy. This evidence suggests that VEGF-A released by podocytes is important for the maintenance of the function and the glomerular structure in the adult kidney. Whether glomerular VEGF-A expression control improves DN has not yet been determined, but there is evidence that shows contradictory results. Administration of anti-VEGF antibodies improved DN in rodents. In experiments conducted in mice, endostatin and tumstatin prevented the development of DN due to a decrease in VEGF-A and angiopoietin 2. In contrast, diabetic mice with gene deletion of VEGF-A in podocytes showed proteinuria and severe diffuse glomerulosclerosis associated with endothelial injury and apoptosis.

The evidence described herein suggests that close monitoring of glomerular VEGF-A levels in diabetes is required in order to avoid adding new lesions or worsening DN. Monitoring glomerular VEGF-A expression within very close margins may have a therapeutic potential, but the optimal concentrations and the right moment to perform such manipulation have not yet been defined.

**VEGF-A relationships with insulin receptors, nephrin and ROS in DN**

In DN, glomeruli with different lesion degrees coexist; VEGF-A expression and its signalling cascade have been related to glomerular changes. In biopsies of patients with DN, there has been evidence of a higher VEGF expression in the glomeruli with lesions due to diabetes than in intact glomeruli. However, VEGF-bound receptor expression was seen to be elevated in glomeruli with mild lesions and decreased in glomeruli with moderate or severe compromise. A similar behaviour was observed with phosphorylation of serine/threonine protein kinase, a protein located in the VEGF signalling cascade, which suggested that other factors would modulate VEGF/VEGFR activity.

Podocytes express insulin receptors, whose activity depends on nephrin expression. Insulin receptors are located in the SD, where podocytes express nephrin and...
VEGFR2\textsuperscript{33,46}. We have characterised the existing interaction between nephrin and VEGFR2\textsuperscript{15}. VEGF overexpression in podocytes was found to decrease nephrin expression and phosphorylation\textsuperscript{16,33}. Hale et al. reported that insulin increases VEGF-A production in podocytes, both in humans and mice\textsuperscript{49}. In transgenic mice, this VEGF-A increase was disrupted by insulin resistance, anticipating the development of podocyte lesions secondary to insulin resistance\textsuperscript{45}. In patients with insulin resistance caused by diabetes and by other diseases, kidney alterations, such as hyperfiltration, proteinuria, modifications in FGB and mesangium were described\textsuperscript{47,48}. Jointly, these findings suggest that VEGF, nephrin and insulin receptor may be related to DN and insulin resistance, thus constituting glomerular pathways susceptible to being modified. Furthermore, oxidative stress secondary to hyperglycaemia may modify glycoalyx, increase ROS and advanced glycation end products, and alter the endothelium. In addition, protein kinase C (PKC) glomerular activation was associated with mesangial expansion, GBM thickening, endothelial dysfunction, cytokine and TGF-Beta activation\textsuperscript{7,15,21,40,41}. Mima et al. described that hyperglycaemia alters nephrin phosphorylation in diabetic rats and cultured podocytes exposed to high concentrations of glucose\textsuperscript{49}. Nephrin phosphorylation interruption was attributed to a “glomerular VEGF resistance” status related to PKC activation\textsuperscript{49}. The VEGF signalling cascade in podocytes and endothelial cells was selectively inhibited by hyperglycaemia\textsuperscript{49}. The increase in glucose and diabetes would cause higher podocyte apoptosis and endothelial dysfunction, partly due to a higher activation of mitogen-activated protein kinase (PKC\textalpha/p38) and SRC homology-2-domain-containing phosphatase-1 (SHP-1) overexpression\textsuperscript{49}. In addition, SHP-1 negatively regulates VEGFR2 and the insulin receptor\textsuperscript{49}. Warren et al. showed that hyperglycaemia reduces endothelial VEGFR2 activity in diabetes\textsuperscript{41}. ROS generation caused by hyperglycaemia was observed to induce VEGFR2 activation and its subsequent breakdown, notwithstanding the VEGF-A\textsuperscript{41}. This would alter the normal response of endothelial cells to circulating VEGF-A due to lower receptor availability. By blocking ROS production with antioxidants, VEGFR2 availability and the lack of endothelial response to VEGF-A caused by hyperglycaemia were reverted\textsuperscript{41}. These results suggest that the increase in VEGF-A present from early stages of DN may be secondary to “VEGF-resistance” of the VEGF-R2 caused by higher receptor breakdown in endothelial cells. Jointly, these publications indicate that, in DN, VEGF overexpression in podocytes may be stimulated in an autocrine and paracrine way by a “VEGF-resistance” state. VEGF-A connections with oxidative stress at glomerular level may represent pathways with therapeutic potential.

### Relationship between angiopoietins and VEGF-A in DN

Angiopoietins, which are growth factors involved in angiogenesis, have been related to DN\textsuperscript{15,36}. Plasma levels of angiopoietin 2 are high in diabetic humans and mice, thus altering the angiopoietin-1/angiopoietin-2 ratio. Diabetic mice with lower angiopoietin 1 levels showed aberrant angiogenesis, hyperfiltration, glomerulomegaly and albuminuria, accompanied by VEGF-A and phosphorylated VEGFR2 overexpression. Alterations caused by reduced angiopoietin 1 were seen to be partially prevented by restoring its expression in podocytes of transgenic mice\textsuperscript{36}. These experiments show the importance of angiopoietins and their relationship with VEGF-A in DN pathophysiology. Modification of protein expression at the glomerular level (by manipulating the cells that produce these proteins) is a therapeutic alternative\textsuperscript{36}.

### Relationship between VEGF-A and nitric oxide in DN

VEGF-A stimulates NO production by means of endothelial NO synthase (eNOS) activation\textsuperscript{15,35,50}. The effects of VEGF-A on vasodilation and on the vascular permeability increase are mediated by the increase in eNOS-dependent NO\textsuperscript{15,17,35,50}. Under normal conditions, VEGF-A induces eNOS activation and an increase in NO; this increase negatively regulates VEGF-A and CTGF, inhibiting ECM build-up\textsuperscript{35}. In diabetes, this relationship changes: the increase in VEGF-A coexists with lower eNOS activity, and there is VEGF-A and NO decoupling\textsuperscript{50}. We showed that VEGF overexpression in podocytes of eNOS KO mice, induced indistinguishable changes of the advanced DN\textsuperscript{35}. In the absence of diabetes, these transgenic mice developed proteinuria, kidney failure and nodular glomerulosclerosis\textsuperscript{35}. This evidence suggests that alterations in glomerular VEGF-A/NO-eNOS relationship are critical and very dangerous, highlighting these events and their relationship with VEGF-A as treatment targets at the glomerular level.

Endothelial NO deficiency secondary to reduced eNOS activity may also associate insulin resistance mechanisms with endothelial dysfunction\textsuperscript{47,48}. Endothelial cells express insulin receptors. By means of eNOS activation, these receptors control vascular tone by inducing vasodilation. For example, in patients with diabetes there are alterations in eNOS activation, establishing a relationship between NO and endothelial insulin resistance\textsuperscript{47-49}. These findings suggest that VEGF-A and the glomerular NO/eNOS ratio may be implied in the insulin resistance status associated with prediabetes, diabetes and CKD.

### Conclusions

Population studies reveal an increasing prevalence of type 2 diabetes worldwide, which suggests that DN will become an even more serious problem. It is imperative to look for alternatives for the diagnosis, prevention and treatment of DN. Going further in the study of molecular pathways with therapeutic potential, such as angiogenic factors, the glomerular VEGF resistance status, insulin resistance in podocytes, the VEGF2/nephrin relationship, VEGF/insulin receptors/nephrin relationship, and the VEGF/NO-eNOS relationship, may provide solutions to the urgent problem of DN in the world.
Key concepts

1. Diabetes mellitus and CKD prevalence have increased in recent decades. The most frequent isolated cause of CKD is DN. Factors related to DN development are: age over 65, uncontrolled hyperglycaemia, hypertension, dyslipidaemia, male gender, smoking habit, family history, and Hispanic or Afro-American origin.

2. Glucose directly and indirectly stimulates VEGF-A cell expression. In DN, there is a systemic and glomerular increase in VEGF-A, but glomerular VEGF-A and the glomerular VEGF-A/NO-eNOS relationship are key factors in DN pathophysiogenesis.

3. Endothelial cells and podocytes express insulin receptors. Nephrin is essential for the action of the insulin receptor in podocytes; its activation is related to VEGF-A. VEGFR2 and nephrin interact in podocytes. Insulin receptors, nephrin and VEGF-A receptors may be mechanistically related to DN and insulin resistance.

4. In DN, VEGF overexpression in podocytes may be stimulated in an autocrine and paracrine way by a “VEGF-resistance” status, in which PKC and ROS would be involved. VEGF-A connections to oxidative stress at the glomerular level may represent pathways with a therapeutic potential for DN.

5. Angiogenic factors, such as VEGF-A and angiopeptins, the relationship of VEGF receptor 2/nephrin, VEGF/insulin receptors/nephrin and the relationship of VEGF/NO-eNOS, VEGF-A/insulin receptors/nephrin, and VEGF/NO-eNOS, represent glomerular pathways that have a crucial significance and may be potential treatment targets for DN.

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