Nitric Oxide Inhibition Induces Early Activation of Type I Collagen Gene in Renal Resistance Vessels and Glomeruli in Transgenic Mice

Role of Endothelin

Christos Chatziantoniou,* Jean-Jacques Boffa,* Raymond Ardaillou,* and Jean-Claude Dussaule†
*Institut National de la Santé et de la Recherche Médicale U.489, Hôpital Tenon, Paris 75020, France; and †AP-HP, Faculté de Médecine St. Antoine, Paris 75012, France

Abstract

Hypertension is often associated with the development of nephroangio- and glomerulo-sclerosis. This pathophysiological process is due to increased extracellular matrix protein, particularly type I collagen, accumulation. This study investigated whether nitric oxide (NO) synthesis is involved in the mechanism(s) regulating activation of the collagen I gene in afferent arterioles and glomeruli. Experiments were performed on transgenic mice harboring the luciferase gene under the control of the collagen I-a2 chain promoter (procolla2(I)). Measurements of luciferase activity provide highly sensitive estimates of collagen I gene activation. NO synthesis was inhibited by Nω-nitro-L-arginine methyl ester (L-NAME) (20 mg/kg per day) for a period of up to 14 wk. Systolic blood pressure was increased after 6 wk of treatment (117 ± 2 versus 129 ± 2 mmHg, P < 0.01) and reached a plateau after 10 wk (around 160 mmHg). Luciferase activity was increased in freshly isolated afferent arterioles and glomeruli as early as week 4 of L-NAME treatment (150 and 200% of baseline, P < 0.01, respectively). The activation of procolla2(I) became more pronounced with time, and at 14 wk increased four- and tenfold compared with controls in afferent arterioles and glomeruli, respectively (P < 0.001). In contrast, luciferase activity remained unchanged in aorta and heart up to 8 wk and was increased thereafter. Increased histochemical staining for extracellular matrix protein deposition, and particularly of collagen I, was detected in afferent arterioles and glomeruli after 10 wk of L-NAME treatment. This fibrogenic process was accompanied by an increased urinary excretion rate of endothelin. In separate experiments, the stimulatory effect of L-NAME on collagen I gene activation was abolished when animals were treated with bosentan, an endothelin receptor antagonist. Similarly, bosentan reduced the increased extracellular matrix deposition in afferent arterioles and glomeruli during NO inhibition. Interestingly, bosentan had no effect on the L-NAME-induced increase of systolic pressure. These data indicate that NO inhibition induces an early activation of the collagen I gene in afferent arterioles and glomeruli. This activation in the kidney precedes the increase in blood pressure and the procolla2(I) activation in heart and aorta, suggesting a specific renal effect of NO blockade on collagen I gene expression that is independent of increased blood pressure and, at least partly, mediated through stimulation of the endothelin receptor. Use of procolla2(I) transgenic mice provides a novel and efficient model to study the pathophysiological mechanism(s) regulating renal fibrosis. (J. Clin. Invest. 1998. 101:2780–2789.) Key words: hypertension • nephroangiosclerosis • renal fibrosis • extracellular matrix

Introduction

Renal vascular and glomerular sclerotic injury is one of the most frequent complications observed in human and experimental hypertension (1). The associated histological lesions include glomerular ischemia, vascular hypertrophy, and interstitial fibrosis. The renal vasculature undergoes structural changes due to extracellular matrix, particularly collagen type I, accumulation (2). This stimulation of extracellular matrix protein synthesis may occur as an adaptation to increased wall tension and/or to the action of different local vasoactive systems.

The endothelium could participate in this pathophysiological process, since it regulates the vascular tone by releasing vasoactive agents such as nitric oxide (NO)1 and endothelin (3, 4). Several in vivo and in vitro studies indicated that NO is an important inhibitor of vascular smooth muscle cell growth (5, 6). On the contrary, endothelin displayed mitogenic properties and induced protein synthesis in cultured vascular smooth muscle cells and mesangial cells (7–9), whereas endothelin antagonism was accompanied by reversal of vascular hypertrophy in the DOCA-salt–, angiotensin II–, or L-NAME–induced hypertension in the rat in vivo (10–12).

The objective of these studies was to evaluate whether or not and by which mechanism(s) NO is involved in the fibrogenic process observed in the renal vasculature during hypertension. For this reason, we have applied the NO-deficiency hypertensive model to transgenic mice, a model extensively studied in the rat (13–15). We used a transgenic mouse line that expresses two reporter genes, luciferase and β-galactosidase, under the control of the promoter of collagen type I gene.

Abbreviations used in this paper: L-NAME, Nω-nitro-L-arginine methyl ester; LU, light units; NO, nitric oxide; procolla2(I), promoter of the α2 chain of mouse collagen type I.

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Because it allows changes in the expression of the collagen I gene to be detected in a highly sensitive manner (16), afferent arterioles and glomeruli were isolated from these mice using a method based on that previously described for the rat (17, 18), and the procol1 gene activation in these structures was compared with that found in heart and aorta. In addition, the role of endothelin in this fibrogenetic process was investigated using pharmacological blockade of endothelin receptors. Our findings suggest that NO plays a major role in the regulation of extracellular matrix synthesis in the renal vasculature. Inhibition of NO synthesis appears to induce vascular and glomerular fibrosis independently of the increase in blood pressure. This physiopathological mechanism is at least partly mediated by an endothelin-induced activation of collagen I gene expression.

Methods

Animal treatment. Male transgenic mice weighing 25–35 g (aged 2–6 mo) at the time of the experiments were maintained on a normal salt diet. Animals had free access to chow and tap water. This transgenic line, named pGB 19.5/13.5, was generated in the laboratory of B. de Crombrugghe (University of Texas, Houston, TX; reference 16). These animals harbor a construction containing the sequences −19.5 to −13.5 kb and −350 to +54 bp of the promoter of the α2 chain of mouse collagen type I [procol1] gene linked to two reporter genes, the firefly luciferase and the Escherichia coli β-galactosidase (see Fig. 1, left and middle panels). The choice of these mice was based on data showing that this construction contains a far-upstream enhancer element regulating high levels of expression of the mouse procol1 gene, and that the expression pattern of the two reporter genes in embryos closely correlates with cell and tissue distribution of collagen I (16). In preliminary experiments, we confirmed these previous findings (see Fig. 1, right panel).

To inhibit NO synthesis, mice were treated with Nω-nitro-L-arginine methyl ester (L-NAME), a NO synthase inhibitor (20 mg/kg per day). In preliminary experiments we found that this dose produced a gradual elevation of blood pressure. In a separate group of control- or L-NAME–treated mice, the mixed endothelin receptor (ET<sub>A</sub> and ET<sub>B</sub>) antagonist bosentan (19) was administered (20 mg/kg per day). Briefly, embryos were fixed for 45 min in a paraformaldehyde solution, and then stained with X-gal (Boehringer Mannheim) overnight at 37°C.

Measurement of blood pressure. Systolic blood pressure was measured by the tail-cuff method adapted to the mouse. A piezoelectric sensor (Sensionor 840–01) connected to a carrier amplifier (Kent 2) was used to detect and convert heart pulses to electric signals. The outputs of the pressure transducer were interfaced to a data acquisition system composed of a Power PC Macintosh 4400/200 computer and a MacLab/4s 16-bit analog to digital converter (ADInstruments, Castle Hill, Australia) allowing sampling at 40,000 samples/s. Pressure recording was analyzed using the Chart module of the MacLab software.

To avoid variations in blood pressure due to day cycle, all measurements were carried out between 9 and 11 A.M. Animals were acclimated for several days before measurements were made. Only animals that did not display signals of stress and that showed stable and reproducible values of blood pressure for at least three consecutive days (> 90% of total cases) were considered for blood pressure measurements. Ten measurements from each mouse were taken at 2-min intervals. The highest and lowest values obtained from each mouse were discarded, and a mean value was determined.

Renal histology. Kidneys from at least three mice from each group were immersed in Dubnoff solution. After fixation, two to three cortical slices of each kidney were embedded in paraffin after conventional processing (alcohol dehydration), and 3 μm–thick sections were stained with Masson trichromic solution for specific staining of extracellular matrix proteins.

Morphologic evaluation. Sections of kidneys were examined on a blinded basis for the level of glomerular sclerosis and microvascular injury using the 0–4+ injury scale, according to established methodology (13). Injury scale 0 means no damaged glomeruli, while 1, 2, 3, and 4+ correspond to 1–25, 26–50, 51–75, and 76–100% of injured glomeruli, respectively. 30–40 samples (containing at least 20 glomeruli per sample) were studied in each group of animals.

Immunohistochemistry. Mice were anesthetized with pentobarbital, and kidneys were removed and immediately fixed in 3% paraformaldehyde/0.1M phosphate buffer, pH 7.4, for 1 h. Renal cortex was cut out into small blocks, infused with 20% sucrose in PBS, and frozen in liquid nitrogen–cooled isopentane. 4 μm–thick cryostat sections were preincubated in PBS containing 1% BSA and 5% decomplemented goat serum, immunostained with an anti-collagen type I (1 mg/ml; Rockland, Gilbertsville, PA) at 1:200 dilution followed by FITC-coupled goat anti-rabbit IgG (5 mg/ml; Organon Tecknika-Capp, Durham, NC), and washed with PBS/0.1% Tween. Antibodies were diluted in PBS containing 0.1% BSA and 0.5% goat serum. Sec-
tions were mounted in an antibleach/glycerol/PBS solution (Citifluor Ltd., London, UK). Photomicrographs were obtained with a Leitz photomicroscope equipped with epifluorescence illumination using Kodak Ectachrome films (Eastman Kodak Co., Rochester, NY).

Measurement of endothelin. Urine samples of mice from the control and the 10th wk L-NAME groups were collected from the bladder of animals. Immunoreactive endothelin-1 was measured by RIA using a Peninsula (Merseyside, UK) commercial kit as described previously (21). Each group included eight mice. Urine endothelin concentration was normalized to urine creatinine concentration, and values were expressed as pg of endothelin per μmol of creatinine.

Acute administration of endothelin. In a separate group of animals, endothelin was injected intraperitoneally (2 nmol/kg), and renal cortical slices were isolated 4 and 20 h after injections. In these renal tissues luciferase activity was measured as described above.

Proteinuria. Urines of 12 mice from each group were collected from the bladder of anesthetized animals just before starting the ice-cold saline perfusion. Samples were centrifuged at 1,500 g for 5 min, and proteins were measured by colorimetry using a specific commercial kit for proteinuria (Red of pyrogallol; Merck, Darmstadt, Germany). Urine protein concentration was normalized using urine creatinine concentration as reference, and values were expressed as milligram of protein per micromol of creatinine.

Statistical methods. Statistical analyses were performed using ANOVA followed by Protected Least Significance Difference Fisher’s test of the Statview software package. Results with $P < 0.05$ were considered statistically significant. All values are means±SEM.

Results

Validation of the model. A primary goal was to investigate whether or not NO is involved in the mechanisms of renal vascular fibrosis observed during the development of hypertension. To this end, an animal model was needed where changes in the activation of extracellular matrix formation could be detected in a sensitive and quantitative way. The procolα2(I) transgenic mouse offers the advantage of high sensitivity of luciferase activity measurements. This advantage, combined with the fact that collagen I is scarcely present in the renal vasculature under normal conditions, made the idea to use the procolα2(I) transgenic line to study renal fibrosis very attractive. However, our preliminary experiments (Fig. 1, right panel) and the previous studies concerning this transgenic line were carried out on embryos or new born animals (16), and several controls were necessary to validate this model in adult animals.

First, we checked whether the luciferase activity in different tissues correlates well with the collagen I distribution in adult animals. Fig. 2A shows the values of luciferase activity in several tissues obtained from 2- to 6-mo-old mice. It is clear that tail and skin displayed high levels of luciferase activity (around 22,000 and 8,000 LU/μg, respectively), aorta showed moderate values (around 1,700 LU/μg), heart, renal cortex, and afferent arterioles had lower values (150, 160, and 200...
Lucretia values were expressed as light units per microgram of protein; 

Table I. Luciferase Levels in Tissues of 2, 3, and 6-mo-old Transgenic Mice

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Age 2</th>
<th>Age 3</th>
<th>Age 6</th>
<th>n</th>
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<tr>
<td>Tail</td>
<td>22108±2159</td>
<td>21198±1376</td>
<td>22147±1079</td>
<td>24</td>
</tr>
<tr>
<td>Skin</td>
<td>8132±783</td>
<td>8256±927</td>
<td>7654±912</td>
<td>24</td>
</tr>
<tr>
<td>Aorta</td>
<td>1492±248</td>
<td>1949±201</td>
<td>1701±189</td>
<td>24</td>
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<td>Heart</td>
<td>134±22</td>
<td>151±19</td>
<td>126±13</td>
<td>24</td>
</tr>
<tr>
<td>Renal cortex</td>
<td>161±11</td>
<td>164±15</td>
<td>193±21</td>
<td>24</td>
</tr>
<tr>
<td>Affer. arterioles</td>
<td>201±10</td>
<td>235±35</td>
<td>205±22</td>
<td>6</td>
</tr>
<tr>
<td>Glomeruli</td>
<td>24±3</td>
<td>21±2</td>
<td>20±2</td>
<td>6</td>
</tr>
</tbody>
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Figure 3. Systolic blood pressure measured in transgenic mice treated with L-NAME for up to 14 wk. Values are means±SEM of 15 mice per group. *P < 0.05 versus control.

Effects of L-NAME treatment. L-NAME, a potent inhibitor of NO-synthase, was given chronically to induce hypertension. After 6 wk of L-NAME treatment, systolic pressure was significantly higher (129±2 versus 116±2 mmHg, P < 0.01, for 6 wk and control, respectively; Fig. 3). Systolic pressure continued to rise with increasing length of L-NAME treatment and reached a plateau around 10–12 wk (156±4 versus 158±3 mmHg, for 10 and 12 wk, respectively; Fig. 3). Staining for extracellular matrix after 6 wk of L-NAME treatment (Fig. 4A) was similar to untreated controls (data not shown). On the contrary, increased levels of extracellular matrix formation were slightly detected after 8 wk and became more apparent after 10 wk in glomeruli and small vessels indicating induction of renal vascular fibrosis (Fig. 4B). Semiquantitative evaluation indicated greater glomerular injury in L-NAME–treated versus control mice (Fig. 5). Glomeruli with injury scores from
2 to 4+ were negligible in controls, whereas they represented one third of the total glomerular population in animals treated with L-NAME for 14 wk. At least a part of the glomerular fibrosis was due to increased presence of collagen I, since immunohistochemistry of cryostat sections indicated increased collagen I positive staining in glomeruli of mice treated for 10 wk with L-NAME (Fig. 6). Proteinuria remained unchanged compared with control values throughout the L-NAME treatment (2.47 ± 0.41 versus 2.71 ± 0.64 and 2.75 ± 0.77 mg/mmol, for control, 8 and 14 wk of L-NAME treatment, respectively). Data obtained after X-gal staining were inconclusive, because renal

Figure 4. Representative example of extracellular matrix staining by Masson trichromic solution in a mouse treated with L-NAME for 6 (A) and 10 wk (B). Note the normal aspect of glomerulus at 6 wk, and the intense green staining (indicating extracellular matrix) and the flocculus retraction of glomerulus at 10 wk. Bar, 10 µm.

Figure 5. Degree of injury in glomeruli isolated from transgenic controls and mice treated with L-NAME or L-NAME + bosentan for 14 wk. *P < 0.05 versus control; #P < 0.05 L-NAME versus L-NAME + bosentan–treated group.

Figure 6. Representative example of immunostaining with collagen I antibody in mouse treated with L-NAME for 10 wk (A) versus vehicle-treated animal (B). Note the intense mesangial (left-pointing arrow), peri-capillary (middle arrow) and interstitial (right arrow) staining.
cortex expresses endogenous β-galactosidase that cross reacts with X-gal. Thus, it was difficult to distinguish the L-NAME–induced from the endogenously produced β-galactosidase (data not shown).

Interestingly, inhibition of NO by L-NAME increased luciferase activity in the renal vasculature earlier than the onset of blood pressure. Thus, isolated glomeruli displayed an almost twofold increase of luciferase activity after 4 wk of treatment (36±2 versus 20±2 LU/μg, P < 0.05, for 4 wk and control, respectively; Fig. 7, top). The L-NAME-induced activation of procolα2(I) gene was further increased with time and reached an eightfold increase compared with controls after 14 wk of treatment (152±13 LU/μg; Fig. 7, top). A similar pattern was observed in afferent arterioles. Luciferase activity started to increase after 4 wk of L-NAME treatment and reached a threefold increase at 14 wk (184±4 versus 252±10 and 591±54 LU/μg, P < 0.05, for control, 4 wk and 14 wk L-NAME treatment; Fig. 7, middle). Procolα2(I) gene activation was detected after 6 wk of treatment in renal cortical slices and continued to increase up to 14 wk (157±6 versus 206±8 and 438±16 LU/μg, P < 0.01, for control, 6 wk and 14 wk L-NAME treatment; Fig. 7, bottom).

To investigate whether this early activation of collagen I gene was specific to the renal vasculature, luciferase activity was measured in the abdominal aorta and the heart of L-NAME–treated mice. Contrary to glomeruli, renal microvessels, and cortex, activation of procolα2(I) gene in aorta was observed much later. 10 wk of NO inhibition were required at least to detect an increase in the expression of procolα2(I) gene in aorta (1,547±107 versus 3,100±145 LU/μg, P < 0.01, for control and 10 wk–treated mice, respectively; Fig. 8, top). The ki-
Nergic pattern of collagen I gene activation in heart was similar to that of aorta. Luciferase activity in heart became higher after 10 wk of L-NAME treatment (137±10 versus 218±10 LU/μg, P < 0.01, for control and 10 wk–treated mice, respectively). To further investigate a possible involvement of endothelin in the fibrogenic process, bosentan, an antagonist of endothelin receptors (ET<sub>A</sub> and ET<sub>B</sub>), was administered in vivo concomitant to L-NAME treatment.

In vivo administration of bosentan had no effect on systolic blood pressure (119±3 versus 117±2 and 116±3 mmHg, in controls and mice treated with bosentan for 8 and 14 wk, respectively). Similarly, bosentan did not change the systolic pressure of L-NAME–treated rats either at 8 wk (144±4 versus 140±4 mmHg), or at 14 wk (161±5 versus 155±5 mmHg), in agreement with the literature for this dose and period of treatment (12, 24).

Bosentan administration to control animals did not affect the luciferase activity in any of the tested tissues (Figs. 9 and 10). In contrast, antagonism of endothelin completely blocked the L-NAME–induced activation of procollagen II gene in glomeruli at 8 and 14 wk (102±11 versus 20±2 LU/μg at 8 wk, P < 0.001, and 152±13 versus 23±2 LU/μg at 14 wk, P < 0.001, for L-NAME− and L-NAME+ bosentan–treated animals, respectively; Fig. 9, top). Similarly, bosentan completely canceled the increase in luciferase activity induced by L-NAME in afferent arterioles and renal cortex up to 14 wk (Fig. 9, middle).
activity measurements provided us with accurate estimates of
tissular (renal vasculature versus heart and aorta) and tempo-
ral (before, during, and after the establishment of hyperten-
sion) activation of procollagen2(I) gene. A novel finding is that
NO inhibition promoted procollagen2(I) expression in the renal
vasculature earlier than the blood pressure increase. In addi-
tion, we advance some clues for understanding the underlying
mechanism(s) since the L-NAME–induced activation of col-
lagen I gene was canceled in the presence of an endothelin re-
ceptor antagonist. Thus, the new information provided by our
study is: (a) NO synthesis participates in the regulation of col-
lagen I gene activation in the vascular tissue in vivo; (b) an
early stimulation of collagen I gene expression occurs in the re-
nal vasculature during NO inhibition and is independent of the
systemic hemodynamics; and (c) at least a part of the increased
collagen I gene expression is mediated by endothelin.

Expression of collagen type I is negligible in renal resis-
tance vessels and glomeruli under normal conditions. In sharp
contrast, collagen I is highly expressed during the physiopatho-
logical process of renal fibrosis, especially nephroangio-
and glomerulo-sclerosis (2, 23). For this reason, expression of col-
lagen I is an excellent diagnostic index of renal vascular fibro-
sis. The line of procollagen2(I) transgenic mouse offers several
specific advantages. It is possible to study mechanisms initi-
ing fibrogenesis since collagen gene activation precedes pro-
tein formation. Luciferase and beta-galactosidase activity were
colocalized in different tissues with collagen I during embry-
onic development, and in new born and adult animals (refer-
ence 16; Fig. 1, right panel; Fig. 2, top), supporting the notion
that these two reporter genes provide accurate estimates of
collagen I gene expression in vivo. Measurement of luciferase
activity is a very sensitive method. For instance, increased
procollagen2(I) activation was detected at 4 wk (Fig. 7) whereas
increased extracellular matrix formation and high expression
of collagen I were detected by conventional morphology and
immunohistochemistry at 10 wk (Figs. 4 and 6). A few micro-
grams of tissue are enough for accurate measurements, the
range of linearity is large (r = 0.98 in the correlation between
beta-galactosidase versus luciferase; Fig. 2, bottom), and control
tissue values were highly reproducible (SE < 10% of mean
within a tissue). In this transgenic line, the use of our technique

Discussion

This study provides new information about the mechanism(s)
involved in the glomerulo- and nephroangio-sclerotic pro-
cesses observed during hypertension. Specifically, our results
highlight the importance of endogenous NO production and of
the balance between NO and endothelin as mediators in the
stimulation of extracellular matrix synthesis, particularly col-
lagen type I. Activation of collagen I gene was measured in
isolated afferent arterioles and glomeruli during pharmacolog-
cal blockade of NO synthesis. Essential to our approach was
the use of transgenic mice expressing luciferase, a reporter
gene, under the control of the promoter of the alpha2 chain of
collagen I gene. The sensitivity and reproducibility of luciferase

and bottom) suggesting a specific action of bosentan on col-
lagen I gene activation in renal vasculature independent of the
rise of blood pressure.

The inhibitory effect of bosentan on L-NAME–induced ac-
tivation of procollagen2(I) gene was also observed in the nonrenal
vascular tissues. Aorta and heart displayed normal levels of
luciferase activity when bosentan was co-administered with
L-NAME at 14 wk (aorta: 4,033±300 versus 1,918±274 LU/
µg, P < 0.001; heart: 316±18 versus 98±10 LU/µg, P < 0.001,
for L-NAME– and L-NAME+ bosentan-treated animals, re-
spectively; Fig. 10).

To verify whether endothelin can acutely induce collagen
type I gene activation in renal tissue of this transgenic strain in
vivo, intraperitoneal injections of endothelin were performed in
a separate group of animals. Exogenous endothelin slightly
increased luciferase activity in renal cortical slices 4 h after the
injections. Luciferase activity was further increased 20 h after
endothelin administration (155±8 versus 211±17 and 456±39
LU/µg, P < 0.01, for 0, 4, and 20 h, respectively).

Bosentan administration protected kidneys from the
L-NAME–induced fibrosis as evidenced by the reduced levels
of extracellular matrix staining in the representative examples
of the L-NAME+ bosentan and the L-NAME group shown in
Fig. 11. Semiquantitative analysis of fibrosis indicated that
L-NAME–induced fibrosis as evidenced by the reduced levels
of extracellular matrix staining in the representative examples
of the L-NAME+ bosentan and the L-NAME group shown in
Fig. 5.

Figure 11. Representative staining of extracellular matrix deposition by Masson trichromic solution in mice treated with L-NAME (A) and
L-NAME + bosentan (B) for 14 wk, respectively. Note that bosentan blunted the intense green staining characteristic of extracellular matrix in
the renal vascular tissue. Bar, 10 µm.
of isolation of renal resistance vessels and glomeruli (17, 18) allowed the study of early mechanisms leading specifically to renal vascular fibrosis. To our knowledge, this is the first study that uses reporter gene expression to investigate mechanisms related to development of hypertension and vascular fibrosis.

Chronic administration of the NO synthase inhibitor L-NAME provides a relatively new experimental model of hypertension (13, 14, 25, 26). The development of hypertension in this model depends on the length and the dose of L-NAME treatment, and is probably due to endothelial dysfunction (25, 26). Renal function is particularly affected during chronic NO inhibition, since L-NAME administration is accompanied by increased afferent and efferent arteriolar and glomerular capillary pressure, and decreased renal blood flow and glomerular filtration rate (13, 26). Prolonged administration of L-NAME in rats induces structural damage of the renal vasculature, such as aneurysm, vascular wall thickening, macrophage invasion and glomerular and interstitial fibrosis (12). In our case, histological lesions were not detectable in mouse kidneys during the initial phase of hypertension (8 wk). They became evident after the week 10, but they remained moderate compared with what has been observed in the rat, probably because of the dose (20 versus 50 mg/kg) and/or species difference. Interestingly, procolla2 gene activation was detected in afferent arterioles and glomeruli much earlier than in heart and aorta (4 versus 10 wk; Figs. 7 and 8). Our finding is in agreement with previous observations indicating that the appearance of renal lesions and structural damage preceded cardiac and aortic fibrosis (27). These data indicate the importance of the NO pathway in the control of renal vascular remodeling.

The most intriguing observation of our studies is that NO inhibition stimulated collagen I gene expression in the renal microcirculation before the increase in blood pressure (4 versus 6 wk; Figs. 3 and 7). Several in vitro studies support the hypothesis that NO can regulate extracellular matrix protein synthesis independently of its vasodilatory action. Stimulation of iNOS by IFNγ or by lipopolysaccharide reduced, while blockade of iNOS by L-NAME–activated collagen and fibronectin synthesis in cultured mesangial cells (28). NO donors displayed cGMP-mediated antimitotic actions in cultured smooth muscle and mesangial cells (6, 29). The L-NAME–induced fibrogenetic effect could be mediated by TGF-β, since L-NAME increased in parallel TGF-β and collagen synthesis in cocultured endothelial-mesangial cells (30). Other studies suggest that during blockade of NO synthase (by L-NAME), the arginase pathway could be activated, thus leading to increased L-proline synthesis, an essential synthesis substrate to collagen formation (31).

Another important observation was that bosentan, an endothelin receptor antagonist, completely canceled the L-NAME–induced activation of collagen I gene in the renal vasculature during (8-wk) and after (14 wk) the establishment of hypertension. A similar effect was observed in the nonrenal tissues (heart and aorta) at 14 wk indicating that the suppressor effect of endothelin antagonist on collagen I gene activation is general and applies to vascular tissue other than renal. The protective action of bosentan did not depend on its effect on systemic hemodynamics, since bosentan did not change blood pressure in controls or L-NAME–treated mice. The absence of a systemic effect during endothelin antagonism is in agreement with previous studies, where bosentan administration did not alter arterial pressure in spontaneously hypertensive and L-NAME–treated rats, or in rats with renal mass reduction (12, 24, 32, 33).

The protective effect of bosentan implies that endothelin is involved in the development of vascular fibrosis. In agreement with this notion, endothelin-1 induced cellular proliferation and increased protein, particularly collagen I, synthesis in cultured coronary smooth muscle cells (34). In our model, acute administration of endothelin activated collagen I gene in kidneys of the transgenic mice. It is possible that NO inhibition stimulated the endothelin system (by increasing the transcriptional rate of the endothelin gene, or by activating the intracellular signaling pathway of the endothelin peptide). The fact that the rate of urinary excretion of endothelin was 2.5-fold higher in the L-NAME–treated mice supports the first mechanism. Elevation of NO production reduced, whereas inhibition of NO synthesis increased, expression of endothelin-1 transcription levels in human endothelial cells (35) further supporting this hypothesis. Similarly, the production of endothelin from aortic vessels was inhibited in presence of NO and potentiated during NO inhibition (36). In addition, antagonism of endothelin receptors blunted the systemic and renal vasoconstrictor effect produced during NO inhibition in anesthetized rats (37). It is also possible that the action of endothelin system became stronger in the absence of the counterbalancing action of NO.

Several recent studies imply that endothelin plays a major role on the mechanisms of nephroangiogenesis and glomerulosclerosis. Thus, antagonism of endothelin receptors delayed the evolution of renal failure (as evidenced by measurements of creatinine and urine protein, and by histological analysis) and increased the survival rate in rats with renal mass reduction (38, 39). Similarly, in the model of murine lupus nephritis, use of an endothelin receptor antagonist improved renal structural damage and reduced extracellular matrix, including collagen I, III, and IV, laminin and proteoglycan, formation (40). More recently, it was observed that transgenic mice overexpressing human endothelin 1 gene developed glomerulosclerosis and interstitial fibrosis (41). It is noteworthy that the appearance of these renal lesions was not accompanied by changes in arterial pressure thus, corroborating the hypothesis that the endothelin-mediated fibrogenic mechanisms are independent of systemic hemodynamics.

In conclusion, the model of transgenic mouse harboring the luciferase reporter gene under the control of collagen I promoter, permitted to investigate early mechanisms in the development of renal fibrosis. NO plays a major role in the mechanisms controlling collagen I gene expression in the renal vasculature, because prolonged inhibition of NO synthesis induced a local renal activation of collagen I gene. This fibrogenic effect is probably independent of systemic hemodynamics and is, at least partly, mediated by the action of endothelin presumably triggered by inhibition of NO synthesis. These data indicate the importance of the balance between endothelial vasoactivators/vasoconstrictors in the physiopathological mechanisms controlling extracellular matrix synthesis during fibrogenesis.

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References


